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## Structure of active human telomerase with telomere shelterin protein TPP1

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### Abstract

Human telomerase is a RNA–protein complex that extends the 3′ end of linear chromosomes by synthesizing multiple copies of the telomeric repeat TTAGGG<sup>1</sup>. Its activity is a determinant of cancer progression, stem cell renewal and cellular aging<sup>2–5</sup>. Telomerase is recruited to telomeres and activated for telomere repeat synthesis by the telomere shelterin protein TPP1<sup>6,7</sup>. Human telomerase has a bilobal structure with a catalytic core ribonuclear protein and a H and ACA box ribonuclear protein<sup>8,9</sup>. Here we report cryo-electron microscopy structures of human telomerase catalytic core of telomerase reverse transcriptase (TERT) and telomerase RNA (TER (also known as hTR)), and of telomerase with the shelterin protein TPP1. TPP1 forms a structured interface with the TERT-unique telomerase essential N-terminal domain (TEN) and the telomerase RAP motif (TRAP) that are unique to TERT, and conformational dynamics of TEN-TRAP are damped upon TPP1 binding, defining the requirements for recruitment and activation. The structures further reveal that the elements of TERT and TER that are involved in template and telomeric DNA handling—including the TEN domain and the TRAP–thumb helix channel—are largely structurally homologous to those in *Tetrahymena* telomerase<sup>10</sup>, and provide unique insights into the mechanism of telomerase activity. The binding site of the telomerase inhibitor BIBR1532<sup>11,12</sup> overlaps a critical interaction between the TER pseudoknot and the TERT thumb domain.

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Numerous mutations leading to telomeropathies<sup>13,14</sup> are located at the TERT–TER and TEN–TRAP–TPP1 interfaces, highlighting the importance of TER–TERT and TPP1 interactions for telomerase activity, recruitment and as drug targets.

Telomerase is a reverse transcriptase that uniquely forms a ribonuclear protein (RNP) with a rapidly evolving non-coding telomerase RNA, that is required for activity, localization and assembly with TERT and other species-specific proteins<sup>15,16</sup>. Telomerase is usually active only in stem cells and germ line cells; its absence in somatic cells limits their proliferative lifespan owing to the shortening of telomeres beyond a critical length<sup>5</sup>. Telomerase reactivation in most cancer cells is essential for their immortal phenotype<sup>3,4</sup>, whereas many disorders of telomere biology, including various cancers, are a consequence of specific mutations in telomerase components<sup>17</sup>. Among these, TERT is the most frequently mutated, followed by TER<sup>18,19</sup>. Cryo-electron microscopy (cryo-EM) structures of human and *Tetrahymena* telomerase have defined the holoenzyme architectures, components and interactions, and provided insights into its mechanism<sup>8–10,20,21</sup>. However, an understanding of the structural basis of diseases of telomerase insufficiency and the development of drugs to inhibit or restore telomerase activity—a long standing goal to treat premature aging and cancer—have been hindered by the lack of detailed knowledge of the atomic level interactions in the human telomerase TERT–TER catalytic core and with TPP1.

TPP1 is one of six shelter in proteins (TPP1, POT1, TRF1, TRF2, RAP1 and TIN2) that associate in various complexes with telomeric DNA<sup>7,22</sup>. TPP1 has a critical role in the regulation of telomere length, as it both recruits and activates telomerase<sup>6</sup>. TPP1 interacts directly with POT1, which binds the single-stranded telomeric DNA 3′ overhang, and with TIN2, which simultaneously binds TRF1 and TRF2 dimers that interact with the double-stranded DNA<sup>22</sup>. TPP1–POT1 switches from telomerase inhibiting by sequestering the 3′ end of telomeric DNA on POT1, to highly activating when TPP1 is bound to telomerase<sup>6,23</sup>. Biochemistry and genetic studies in cells have shown that the TPP1 oligonucleotide and oligosaccharide-binding fold domain (OB) interacts with TERT through a glutamate- and leucine-rich patch (TEL patch) and the N terminus of the OB domain<sup>24–27</sup> (NOB). POT1 binds TPP1 on a POT1-binding domain separated by a linker that follows the OB domain<sup>28,29</sup> (Fig. 1a). High-resolution cryo-EM structures of *Tetrahymena* telomerase established that p50, a constitutive component that is an orthologue of the activating functions of TPP1, binds TEN and TRAP<sup>10,20</sup>. However, TPP1 interacts only transiently with human telomerase in a cell cycle-dependent manner<sup>30</sup>, making structural studies challenging. Here we purified human telomerase in complex with TPP1 and telomeric DNA and determined their cryo-EM structures. These structures define the determinants of telomerase recruitment to telomeres by TPP1, reveal the unique TERT–TER interactions that regulate telomerase activity, assembly and template–DNA handling, and localize many mutations that lead to diseases of telomerase insufficiency to these interfaces.

## Overall structure

We purified human telomerase with DNA primer d(T<sub>12</sub>TTAGGG) in three steps, as described in Methods. The purified DNA-bound telomerase was incubated with

approximately tenfold excess of the TPP1 OB domain (hereafter referred to as TPP1) (Fig. 1a) before vitrification on cryo-EM grids. As previously described<sup>8,9</sup>, telomerase has a bipartite structure with a catalytic core RNP (Fig. 1b–d) and a H and ACA box (H/ACA) RNP (Fig. 1f, g) flexibly connected by P1a and P4.1–P4.2–P5 regions of TER (Fig. 1e). Separate focused refinements of the two RNPs resulted in resolutions of 3.3–3.8 Å for the TERT–TER catalytic core and 3.2 Å for H/ACA RNP (Fig. 1b, c, f, Extended Data Figs. 1–3). The telomerase H/ACA RNP has a non-canonical function in telomerase biogenesis and localization, rather than pseudouridylation of specific substrates<sup>31</sup>. The H/ACA RNP comprises two complete sets of H/ACA proteins dyskerin, GAR1, NOP10 and NHP2 bound to successive irregular stem-loops (P4 and P7) followed by box H (AGAGGAA) and box ACA nucleotides, respectively, and a single TCAB1 bound at the P8 apical loop with dyskerin and NHP2 (Fig. 1e–g, Extended Data Fig. 4a). The H/ACA RNP model is almost the same as the previously reported one<sup>8</sup> and is therefore not discussed further in this Article.

The catalytic core comprises<sup>32</sup> the TERT and TER template–pseudoknot (t/PK) and conserved regions 4–5 (CR4/5) (Fig. 1e, Extended Data Fig. 4). The conserved RNA-binding domain (RBD), reverse transcriptase (RT) and C-terminal extension (CTE) of TERT form a ring, and the TERT-unique TEN and TRAP (a large insertion in RT fingers) form a complex above the TERT ring<sup>10,20</sup> (Fig. 1a, b, Extended Data Fig. 5a, b). TPP1 binds to TERT TEN and TRAP (Fig. 1a, c, d). CR4/5 extends from P4, and P5, P6 and P6.1 are connected by a three-way junction (Fig. 1e, Extended Data Fig. 4a). About 22% of particles also contain density corresponding to the histone H2A–H2B dimer—which was previously observed at somewhat higher percentage—bound at and possibly stabilizing this three-way junction<sup>8</sup> (Fig. 1d, Extended Data Fig. 3a, b). The overall higher resolution of the catalytic core in the present study enabled us to assign sequence more accurately to cryo-EM density and position side chains (Fig. 1d, Extended Data Fig. 2).

## TPP1 interactions with TEN—TRAP

A notable difference between the structure of the catalytic core without<sup>8</sup> and with TPP1 is that TEN is at a higher resolution in the complex with TPP1 (Fig. 2a). This enabled us to accurately model TEN—previous modelling<sup>8</sup> had around one-third of the residues in the wrong register. The overall structures of human TEN and TRAP, as well as the extended  $\beta$ -sheet between TEN and TRAP, are remarkably similar to those in *Tetrahymena* telomerase (Fig. 2b, Extended Data Fig. 5b). However, the position of TEN–TRAP relative to the TERT ring is different, which may be caused by the double-stranded end of the pseudoknot (P2a.1) abutting TEN in human telomerase (Extended Data Fig. 5c–f). TRAP is connected at its N- and C-terminal ends to the conserved helices IFD<sub>a</sub> and IFD<sub>c</sub>, respectively, by flexible linkers that allow positional dynamics of TEN–TRAP relative to the TERT ring (Fig. 2b).

The architecture of TPP1 bound to telomerase is the same as for the free protein<sup>33</sup>, except at its interface with TEN–TRAP, where loop  $\alpha$ 2– $\beta$ 4 forms the short helix  $\alpha$ 2.1 (Fig. 2c). TPP1 loop  $\alpha$ 2– $\beta$ 4 and NOB<sup>27</sup> are shifted to positions similar to those in *Tetrahymena* telomerase-bound p50<sup>10</sup>, but adopt different conformations (Extended Data Fig. 6a). Comparison of the cryo-EM density of TEN–TRAP in telomerase without and with TPP1 provides insight into

the conformational changes caused by TPP1 binding. TRAP is largely unchanged, whereas the TEN interface with TPP1 and TRAP becomes structured, as evidenced by the stronger density—especially for its  $\beta 3$ – $\beta 4$ ,  $\beta 4$ – $\alpha 6$  linker and  $\alpha 7$  (Extended Data Fig. 6i,j). Thus, TPP1 binding stabilizes the interaction between TEN and TRAP. TPP1 binding also damps the positional dynamics of TEN–TRAP, as evidenced by the improvement in cryo-EM density on the outside surface of TEN (Fig. 2a, Extended Data Fig. 6i, j).

Superposition of TPP1–TEN–TRAP on *Tetrahymena* p50–TEN–TRAP<sup>10</sup> reveals that the binding interfaces are globally the same, including their three-way interaction<sup>20</sup> (Extended Data Fig. 6), but most specific interactions are not conserved (Supplementary Fig. 2). There is a continuous interface between TPP1 and TEN–TRAP (Fig. 2e, Extended Data Fig. 6e) that on TRAP includes interactions with residues previously identified as the TEL patch<sup>24</sup> and NOB<sup>27</sup> (Fig. 2c). We identified three charge–charge interactions, TEN K78–TPP1 E215, previously identified in a charge-swap experiment<sup>34</sup>, TRAP R774–TPP1 E168 (the TEL patch) and TEN D129–TPP1 R92 (NOB) (Fig. 2d). The rest of the interface is primarily hydrophobic, involving mostly leucines on TRAP and TPP1 (Fig. 2f) and short side chains (alanine, glycine and serine) on TEN (Fig. 2e). Among these, TERT mutations R774L and Y772C are associated with dyskeratosis congenita and aplastic anaemia, respectively<sup>13</sup>. Except for K78, most of the residue substitutions or disease-mutation substitutions previously inferred to disrupt direct TPP1–TERT interactions<sup>8,10,35,36</sup> act indirectly, either by destabilizing the TEN–TRAP interface or changing TEN or TRAP structure (Extended Data Fig. 7a–c).

We tested alanine substitutions on TEN–TRAP interface residues using a telomerase direct activity assay to evaluate their importance in TPP1 binding (Methods). TPP1 enhanced telomerase activity approximately twofold, with only a small effect on repeat addition processivity<sup>24</sup> (RAP), similar to *Tetrahymena* p50<sup>37</sup>, under optimized conditions of excess TPP1. All mutants showed significant decreases of this activity stimulation, from approximately 50–85% (Methods), including the well-studied K78A<sup>34</sup> (75% decrease) (Fig. 2g, Extended Data Fig. 7d–f). TRAP L766A, Y772A and L798A caused more than 75% loss of activity stimulation, and R774A (similar to its partner TPP1 E168A<sup>24</sup>) and TEN D129A caused approximately 60% loss of activity stimulation. We next tested whether TPP1 binding defects would affect RAP enhancement by TPP1–POT1. Addition of TPP1–POT1 had little effect on activity but enhanced RAP by approximately 1.4-fold (Extended Data Fig. 7g–i). The TEN–TRAP residue substitutions caused a similar pattern in decreasing stimulation of RAP by TPP1–POT1 to the decreasing stimulation of activity by TPP1 alone (Fig. 2g, Extended Data Fig. 7i). These results validate the identified interface interactions for TPP1 binding. This atomic model of the TEN–TRAP interface for TPP1 binding defines the structural requirements for telomerase recruitment and provides a rational basis for drug design to inhibit telomerase action by repressing recruitment.

## Features of the catalytic cavity

The TER t/PK forms an oval around the TERT ring that is closed by P1b (Fig. 3a, b). The single-strand region of the t/PK that includes the template wraps along the RBD (5′), RT (template) and CTE (3′), where it passes under the TRAP–TEN (Fig. 3a). In

the structure, only the last four bases in the DNA primer d(T<sub>12</sub>TTAGGG) pair with the template, with the duplex shifted out of the active site as previously reported<sup>8</sup> (Fig. 3b). This observation is consistent with duplexes of four (or five if the active site is occupied) base pairs in *Tetrahymena* telomerase structures at multiple steps of telomere repeat synthesis<sup>10</sup>, indicating that a short unstable duplex is a common feature for telomerase<sup>38</sup>.

During telomere repeat synthesis, the template movement boundary is determined by the template boundary element (TBE) acting as an anchor on its 5' side<sup>10,21,39</sup>. In our structure, the perpendicularly positioned P1b-P3 is anchored on TERT RBD by interactions to the single nucleotide junction (C186) and base pairs on either side (Fig. 3c, Extended Data Fig. 4d). This establishes P1b-P3 as the TBE, consistent with mutagenesis results<sup>40</sup>. The template is connected to the TBE by a linker (TBE<sub>L</sub>) of eight nucleotides (U38-U-U-U-U-U-G-U45), versus two in *Tetrahymena* telomerase (Fig. 3b, Extended Data Fig. 8a, b). Although residues U38 to U43 are loosely tethered, as indicated by their weak cryo-EM densities (Extended Data Fig. 8a), G44 and U45 are inserted into a binding pocket on the RBD, which appears to be a secondary 'kedge' anchor (Fig. 3a, Extended Data Fig. 8c-e). As the template moves, nucleotides in the pocket could pop out and be replaced sequentially by the subsequent residues, up to U41-U42 when the TBE<sub>L</sub> would be fully stretched (Extended Data Fig. 8f). The proposed kedge anchor would help keep the template correctly positioned in the catalytic cavity.

The template-DNA duplex is held in the catalytic cavity by template interactions with motif 3 and fingers on RT and DNA interactions with the thumb loop, the thumb helix on CTE and the T motif on the RBD<sup>8</sup> (Fig. 3d, e, Extended Data Fig. 8g). A bridge loop on the RBD, first identified in *Tetrahymena* telomerase and proposed to regulate duplex length<sup>10</sup>, bridges the two ends of the duplex with its conserved tip residues (K499 and H500; R413 and F414 in *Tetrahymena*) inserted on the major groove side (Fig. 3d, e). Individual substitutions of these residues to their *Tetrahymena* counterparts had little effect on activity, whereas substitution with alanine substantially decreased activity and RAP (Extended Data Fig. 8h). K499, together with Y333, E565 (T motif) and R622 (fingers motif), forms a conserved interaction network that regulates stepwise flipping of template bases into the active site (Extended Data Fig. 8i). On the other end of the duplex, the first unpaired DNA nucleotide (dT14) inserts between template nucleotides at the duplex-single-strand junction and interacts with L980 (Fig. 3e). This leucine was proposed to have a major role in strand separation in a recent publication on human telomerase structure<sup>41</sup>. The following T13 flips out and stacks on bridge loop residue H500 (Fig. 3e, Extended Data Fig. 8j). By contrast, in *Tetrahymena* telomerase, all the unpaired DNA nucleotides are stacked continuously under F414 from the bridge loop<sup>10</sup>. This difference may be owing to flexibility of the exiting DNA in the absence of telomeric DNA handling protein POT1 versus the constitutively assembled orthologue Teb1 in *Tetrahymena*<sup>21</sup>. Consistent with this, there is no traceable cryo-EM density for DNA nucleotides beyond T13.

On the template 3' side, the template alignment region enters a narrow, positively charged channel between the TRAP and CTE thumb helix<sup>10</sup> (TRAP-TH channel) and the single-stranded RNA between the template and P2a.1 makes a gooseneck turn below on the CTE (Fig. 3f). The arginines lining the channel could function as a ratchet for template movement

during telomere repeat synthesis, as proposed for *Tetrahymena* telomerase<sup>10</sup>. TPP1 binding damps the dynamics of TEN–TRAP, which we propose helps to form the optimal TRAP–TH channel for regulation of telomeric DNA synthesis. Overall, the common features of the human telomerase catalytic cavity to those in *Tetrahymena* suggest a largely conserved mechanism for repeat synthesis and translocation.

## TERT–TER interactions scaffold the catalytic core

On the 3′ side of the template, the structurally conserved pseudoknot forms an irregular helix that wraps around TERT ring from CTE to RBD (Fig. 4a, b). The pseudoknot can be divided into three regions: P2a.1, which connects to the single strand 3′ of the template<sup>32</sup>; P2a–P2b, which contains a 5-nucleotide (nt) asymmetric internal loop (J2a/b); and P2b–P3, which contains conserved base triples from loop-stem interactions<sup>42,43</sup> (Fig. 4a, b, Extended Data Fig. 4a, b). The tertiary structure of P2b–P3 is almost identical to the structure determined for this sub-domain in solution<sup>43</sup>. Adjacent to P2b–P3 is an approximately 90° bend at the 5-ntJ2a/b bulge between P2b and P2a<sup>44</sup>; the rest of the P2 stem (P2a.1–P2a–P2b) that passes over the CTE is mostly helical (Fig. 4a, b). The interactions of the pseudoknot with the TERT ring are located mainly on P3 (Extended Data Fig. 4d), which fits into the positively charged cleft between the CTE and RBD (Fig. 4a, Extended Data Fig. 9a). Notably, except for a flipped-out G73 between P2a.1–P2a and C116 at the P2b–P3 junction, the entire P2 only contacts TERT at its 5′ end, where P2a.1 abuts a positively charged surface of TEN on its major groove side (Fig. 4b, Extended Data Fig. 9a). Whereas P2b–P3 is fixed on TERT, the rest of the pseudoknot exhibits conformational dynamics, as revealed by modelling of different class averages (Extended Data Fig. 9b–e).

Besides the interactions from P3, the TERT ring is further stabilized by CR4/5 binding from the opposite side. In CR4/5, P6 is stacked on P5.1—a predicted 3-bp helix<sup>36,45</sup> formed by interactions in the large internal loop between P6, P5 and P6.1—and the P6.1 hairpin is perpendicular to P6 (Fig. 4c, Extended Data Fig. 4a, c, e). The highly conserved P6.1 hairpin bridges the RBD and CTE, whereas P6 binds the RBD, as observed in teleost fish<sup>46</sup> (Fig. 4c). Together, CR4/5 and the pseudoknot form a framework that inhibits large-scale conformational change in the TERT ring (Fig. 4b).

A single bulge nucleotide (U177) below the three U–A–U triples<sup>42,43</sup> in P2b–P3 is within about 7 Å of U307 on the P6.1 hairpin loop in CR4/5 (Fig. 4d, Extended Data Fig. 4d, e). Our model reveals that these two nucleotides bind in separate pockets on opposite sides of the CTE, separated by loop L<sub>α32–α33</sub> (amino acids 1019–1028). U177 interacts with four L<sub>α32–α33</sub> residues, including L1019, whereas U<sub>307</sub> hydrogen bonds with Q1023 (Fig. 4d, Extended Data Fig. 4d, e). Disease mutations<sup>13</sup> L1019F (pulmonary fibrosis), V1025F and V1090M (aplastic anemia), and N1028H (DKC) would disrupt interactions that stabilize P2b–P3 and/or P6.1 binding to CTE in this region. Docking of telomerase-specific inhibitor BIBR1532 onto its binding site on CTE, based on homology to CTE of *Tribolium* TERT-like protein<sup>11</sup>, reveals that its benzoic ring would occlude U177 binding and its naphthalene group repositions L<sub>α32–α33</sub>, which could also affect P6.1 interaction (Fig. 4d). Thus, as a non-nucleosidic telomerase inhibitor<sup>12</sup>, BIBR1532 disrupts critical TERT–TER interactions, making it the first known example of a drug targeting a non-template region of TER<sup>47</sup>.

## Insights into catalytic core disease mutations

More than 185 and 75 mutations in TERT and TER, respectively, lead to short telomeres and manifest as diseases of telomerase insufficiency<sup>13</sup>. In TER, the vast majority of these are in the pseudoknot and CR4/5, and there are only a few in the H/ACA RNA<sup>13</sup>. We mapped known disease-associated mutations of TER and TERT (that would not obviously disrupt TERT hydrophobic core) onto the structure (Fig. 5, centre). Almost all TER mutations in the catalytic core contact or are adjacent to nucleotides that contact TERT. Notably, there are many mutations in pseudoknot P3 and P1b (Fig. 5, Extended Data Fig. 4d, e). A few others would affect critical folds such as the conserved 90° bend in the J2a/b loop<sup>44</sup> by disruption of stem base pairs (Extended Data Fig. 4a). The 3.3 Å model of the catalytic core reveals specific TERT-TER contacts that would be affected by individual mutations (Fig. 5, boxed examples). For pulmonary fibrosis, examples include mutations in TERT RT fingers (R622C) and TRAP (R756L) motifs that would disrupt interactions with the template. Fingers R622 is part of the conserved network of residues that regulates sequential flipping of the template nucleotides opposite the active site<sup>8,10</sup> (Fig. 3e). R756 is at the top of the TRAP-TH channel where it stacks with template (A55) and helps regulate the one-way movement of the template during nucleotide addition (Fig. 3f). CTE mutation R1086C would disrupt interactions with the backbone of P2b C116, the last residue of P2b–P3 pseudoknot to interact with TERT (Extended Data Fig. 4d). TER mutations in dyskeratosis congenita include G73U; G73 is the single residue at the P2a.1–J2a/b junction that specifically interacts with CTE (Fig. 4b). Mutations throughout the pseudoknot P3 that have direct contacts with TERT or stabilize the extended P3 stem (Extended Data Fig. 4d), are associated with aplastic anaemia and other diseases<sup>13</sup>. Aplastic anaemia TER mutation G305U would disrupt the top of P6.1 and interaction with CTE Q1023, and TH R979Y would affect template handling in the TRAP-TH channel (Fig. 5). Acute myeloid leukemia G309U (P6.1) and myelodysplastic syndrome R535H (RBD) mutations provide additional examples of mutations that disrupt P6.1–TERT interactions with CTE Q1023 and P6.1 A302, respectively. Dyskeratosis congenita R381P (RBD) would disrupt arginine interactions with three base pairs flanking the bulged U<sub>291</sub> in CR4/5 helix P6. Last, K570N (in the RBD T motif) is associated with Hoyeraal–Hreidarsson syndrome (Fig. 5) and pulmonary fibrosis; K570 interacts with the DNA backbone in the template–DNA duplex (Fig. 3e). Analysis of these interactions validates the importance for telomerase activity of the TERT–TER interactions defined in our model and provides a structural basis for drug targeting.

## Summary

This structure of human telomerase in complex with shelterin protein TPP1 illustrates the detailed interface essential for recruitment and activation. Our comparison of common and divergent features of the catalytic cores of *Tetrahymena*<sup>10</sup> and human telomerase at 3.3 Å resolution has unveiled details of template–DNA handling, template boundary definition and duplex length during telomere repeat addition. TPP1 binding stabilizes the TRAP–TEN interface and TRAP–TH channel, and its association with POT1 positions it to bind the newly synthesized telomeric DNA. Analysis of disease mutations as well as the binding site for the telomerase-specific inhibitor BIBR1532<sup>11,12</sup> revealed the critical importance



of TERT–TER interactions for telomerase activity and as potential drug targets. Overall, these findings provide a basis for structure-guided telomerase-targeted drug discovery and a deeper understanding of disease mutations leading to telomere biology disorders.

## Methods

### Telomerase expression and purification

Human telomerase was reconstituted *in vivo* and purified largely as previously reported<sup>9</sup>, except for the addition of an initial IgG affinity purification step. In brief, human TERT and TER genes from plasmid pBABEpuroUThTERT+U3-hTR-500<sup>48</sup> (AddGene) were individually cloned into pcDNA3.1 vectors, and pcDNA3.1-ZZ-TEV-SS-TERT and pcDNA3.1-U3-TER-500ex-HDV were constructed as described<sup>9</sup>. For the TERT plasmid, a tobacco etch virus (TEV) protease cleavage site was inserted between the tandem protein A (ZZ) and Twin Strep (SS) tag (IBA Lifesciences) cassettes for telomerase purification. For the TER plasmid, the U3 snoRNA promoter<sup>49</sup> was inserted before TER and a 500-nt extended sequence<sup>50</sup> was inserted between TER and hepatitis delta virus ribozyme<sup>51</sup> (HDV). We used adherent HEK 293T cells for telomerase reconstitution. The cells were cultured on 150-mm plates with DMEM media (Gibco) supplemented with 10% FBS (Atalanta), 1% penicillin-streptomycin and 1% GlutaMAX. The cells were transfected with an optimal ratio of 1:4 of TERT and TER plasmids using calcium phosphate reagents<sup>52</sup>. After 48 h of transfection, the cells were harvested by centrifugation (1000x g, 5 min) and resuspended in pre-chilled hypotonic lysis buffer (HLB) (20 mM HEPES·NaOH pH 8.0, 10% glycerol, 2 mM EDTA, 0.2 mM EGTA, and 1 mM TCEP) supplemented with 0.1% NP-40, 0.2 mM PMSF, 10 μM MG-132 (Sigma), and 0.2% protease inhibitor cocktail (Sigma). Cells were incubated on ice for 10 min before being flash frozen and stored at –80 °C.

For telomerase purification, frozen cells from 120 plates were thawed and subjected to two additional freeze-thaw cycles. The cell lysate was adjusted to 300 mM NaCl concentration and incubated on ice for 30 min. An equivalent volume of HLB buffer supplemented with 0.1% Igepal CA-630 was added to the cell extract to dilute the NaCl concentration to 150 mM. The cell lysate was cleared by ultracentrifugation (125,000g) for 1h at 4°C. The prepared cell lysate was incubated with IgG resin (Sigma) for 5 h at 4°C with end-over-end rotation. The resin was washed three times with wash buffer (20 mM HEPES·NaOH pH 8.0, 150 mM NaCl, 10% glycerol, 2 mM EDTA, 0.2 mM EGTA, 1 mM TCEP, and 0.1% Igepal CA-630) before elution by TEV protease cleavage at 4 °C for 1 h. The eluate was subsequently incubated with streptavidin agarose resin (Sigma) prebound to a 5′-biotinylated oligonucleotide<sup>53</sup> at room temperature for 2.5 h. The resin was washed with wash buffer and eluted with 50 μM competitor oligonucleotide<sup>53</sup> in 500 μl wash buffer 3 times. The total elution was further incubated with pre-blocked MagStrep XT3 resin (IBA LifeSciences) overnight at 4 °C along with 5 μM DNA primer, T<sub>12</sub>TTAGGG. The resin was washed with wash buffer and eluted with 35 μl biotin elution buffer (20 mM HEPES·NaOH pH 8.0, 150 mM NaCl, 2 mM MgCl<sub>2</sub>, 20 mM biotin, 1 mM TCEP, and 0.05% Igepal CA-630) to get the final product.

### TPP1 expression and purification

TPP1 OB (residues 88–249) and TPP1 OB-PBD (residues 88–334, Addgene) were cloned into a modified pETduet vector with an N-terminal 6xHis-MBP tag followed by a TEV protease cleavage site and a pET-SUMO vector individually. The proteins were expressed in *Escherichia coli* BL21 (DE3) cells as described<sup>33</sup>. LB media with 100  $\mu\text{g ml}^{-1}$  Ampicillin (for TPP1 OB) or 50  $\mu\text{g ml}^{-1}$  kanamycin (for TPP1 OB-PBD) were used for bacterial growth at 37 °C to an OD<sub>600</sub> of 0.6. Protein expression was induced by 0.5 mM IPTG for 20 h at 18 °C. Cells were harvested, suspended in buffer A (20 mM HEPES-NaOH pH 8.0, 1 M NaCl, 15 mM imidazole, 5% glycerol, and 1mM TCEP), sonicated and centrifuged at 34,000g for 45 min. The supernatant was loaded onto a Ni-Sepharose affinity column (HisTrap HP; GE Healthcare). The column was washed with 95% buffer A and 5% buffer B (20 mM HEPES-NaOH pH 8.0, 0.5 M NaCl, 1 M imidazole, 5% glycerol, and 1 mM TCEP), and then the protein was eluted with 70% buffer A and 30% buffer B. The eluate was incubated with TEV protease (for TPP1 OB) or SUMO protease (for TPP1 OB-PBD) for 3–4 h at room temperature while dialyzing against buffer containing 20 mM Tris-HCl pH 7.5, 250 mM NaCl, and 5 mM  $\beta$ -mercaptoethanol. TPP1 was collected from the flow through after loading the cleavage reaction onto a Ni-Sepharose affinity column. Further purification was conducted by size-exclusion chromatography (SEC) (HiLoad 26/600 Superdex 75; GE Healthcare) in SEC buffer (20 mM HEPES-NaOH pH 8.0, 150 mM NaCl, 5% glycerol, and 1 mM TCEP). TPP1 OB protein peak fractions were pooled and concentrated to 300  $\mu\text{M}$  using 10 kDa cut-off Amicon filters (Millipore Sigma), then aliquoted and stored at –80 °C. To prepare the TPP1 bound telomerase, a final concentration of 500 nM purified TPP1 OB was added to the purified telomerase and incubated on ice for 1 h just prior to grid preparation.

### TPP1–POT1 complex purification

POT1 was expressed using the Bac-to-Bac system (Thermo Fisher Scientific) in Sf9 cells. In brief, cDNAs encoding POT1 was chemically synthesized by IDT (Integrated DNA Technologies) and cloned into pFastBac1 vector (Thermo Fisher Scientific) with an N-terminal 6xHis-MBP tag followed by a TEV protease cleavage site. The expression vectors were used to make baculoviruses based on the established protocol for Bac-to-Bac system (Thermo Fisher Scientific). Sf9 cells ( $2.0 \times 10^6 \text{ ml}^{-1}$ ) were transfected with viruses using a multiplicity of infection (MOI) of 3 at 27 °C in SF-900 II SFM media (Thermo Fisher Scientific). The cells were harvested 48 h after infection, and proteins were purified as described for TPP1. Fractions of POT1 collected from SEC (HiLoad 26/600 Superdex 200; GE Healthcare) were mixed with TPP1 OB-PBD at 1:1 ratio and incubated at 4 °C for 1 h before loaded to another SEC. TPP1-POT1 fractions were collected and concentrated to around 100  $\mu\text{M}$  using 30 KDa cut-off Amicon filters (Millipore Sigma), then aliquoted and stored at –80 °C.

### Cryo-EM sample preparation and data collection

Telomerase samples were applied to glow-discharged lacey gold grids coated with a supporting ultrathin carbon film (EMS). The grids were blotted with Grade 595 filter paper (Ted Pella) and flash-frozen in liquid ethane using an FEI Vitrobot Mark IV. Cryo-

EM grids prepared and frozen with varying parameters (including the time and gas for glow-discharging, the volume of applied sample, temperature and humidity of the Vitrobot sample chamber, waiting time before blotting, blotting time and force) were screened in an FEI Tecnai TF20 transmission electron microscope for optimal ice thickness, particle concentration and particle distribution on grids. The best grids were obtained with 30 s glow-discharging in mixed H<sub>2</sub> and O<sub>2</sub>, 5 µl sample applied on each grid, and Vitrobot set at 10°C, 100% humidity, waiting time 4 min, blotting force 4, and blotting time 10 s.

Cryo-EM grids were loaded into a ThermoFisher Titan Krios electron microscope operated at 300 kV for automated data collection using SerialEM<sup>54</sup>. Movies of dose-fractionated frames were acquired with a Gatan K3 direct electron detector in super-resolution mode at a pixel size of 0.55 Å on the sample level. A Gatan Imaging Filter (GIF) was installed between the electron microscope and the K3 camera with the slit width setting to 20 eV. The microscope was carefully aligned prior to each imaging session and the parallel beam was optimized using coma-free alignment in SerialEM<sup>54</sup>. The dose rate on the sample was set to approximately 55 eT Å<sup>-2</sup> in total, which was fractionated into 50 frames with 0.06 s exposure time for each frame. Finally, 31,728 movies were collected in three separate imaging sessions with three batches of cryo-EM grids (Extended Data Fig. 1).

### Cryo-EM data processing

The cryo-EM data processing workflow is outlined in Extended Data Fig. 1. All steps described below were performed with RELION 3.1<sup>55</sup> unless otherwise indicated. Dose-fractionated frames of each movie were 2× binned (pixel size of 1.1 Å), aligned for the correction of beam-induced drift, and dose weighted using RELION's implementation of UCSF MotionCor2<sup>56</sup>. Contrast transfer function (CTF) parameters, including defocus and astigmatism, of each dose-weighted micrograph were determined by CTFFIND4<sup>57</sup> within RELION.

Three datasets, one for each data collection session, were initially processed separately in three batches (Extended Data Fig. 1a). For dataset1, 666,560 particles were picked from 2,000 representative micrographs using template-free auto-picking in RELION. These particles were extracted in boxes 384 pixels square and 3× binned to 128 pixels square (pixel size of 3.3 Å) for 2D classification. 203,042 particles from the best classes were selected to train a particle detection model in Topaz<sup>58</sup> for subsequent neural-network based particle picking. From the 16,244 micrographs of dataset1 a total of 8,500,796 particles were picked, 3x binned and extracted in boxes 128 pixels square (pixel size of 3.3 Å). Two rounds of 2D classification were performed to remove 'bad' particles in classes with fuzzy or uninterpretable densities. 3,006,138 particles in classes with clear features for both the catalytic core and H/ACA RNP were selected. Refinement of the selected particles using a previously published cryo-EM map<sup>9</sup> (EMD-7521) as an initial model generated a 3D reconstruction with only low-resolution features. To improve the resolution, each particle was re-extracted into two sub-particles with shifted origins (RELION option: --recenter\_x/y/z), one centred on the catalytic core and the other centred on the H/ACA RNP. Sub-particles centred on the catalytic core were extracted in boxes 128 pixels square (pixel size of 2.2 Å) and refined using a soft mask encasing the catalytic core (mask1). The

resulting orientation parameters for each sub-particle were used as inputs for the subsequent masked 3D classification step with local angular search (RELION options: --sigma\_ang 30 --healpix\_order 2). In parallel, sub-particles centred on the H/ACA RNP were extracted in boxes 200 pixels square (pixel size of 2.2 Å), refined and 3D classified with a soft mask covering the H/ACA RNP (mask2). An additional step of 2D classification was performed before 3D refinement to remove sub-particles that were not centred properly on the H/ACA RNP. Last, 1,781,404 sub-particles centred on the catalytic core and 1,055,677 sub-particles centred on the H/ACA RNP were selected from dataset1.

Dataset2 and dataset3 were processed in a similar way as described above for dataset1 (Extended Data Fig. 1a). For dataset2, 4,413,286 particles were picked from 8,361 micrographs. After screening, 429,103 sub-particles centred on the catalytic core and 439,516 sub-particles centred on the H/ACA RNP were selected. For dataset3, 2,641,248 particles were picked from 7,123 micrographs. 403,764 sub-particles centred on the catalytic core and 504,954 sub-particles centred on the H/ACA RNP were selected after screening. The two types of sub-particles from the three datasets were combined and processed respectively with the following steps.

For the H/ACA RNP, the combined 2,000,147 sub-particles (pixel size of 2.2 Å) were refined to 4.4 Å resolution using mask2 (Extended Data Fig. 1c). Then, two rounds of 3D classification were performed with local angular search in 30° (RELION options: --sigma\_ang 30 --healpix\_order 2) and 12° (RELION options: --sigma\_ang 12 --healpix\_order 3), respectively. 432,108 sub-particles from the best class were extracted in boxes 320 pixels square (pixel size of 1.1 Å) and refined to 3.3 Å resolution. After another round of 3D classification with a finer angular sampling interval (RELION options: --sigma\_ang 8 --healpix\_order 4), 256,859 sub-particles were selected for 3D refinement, following by CTF refinement to correct CTF parameters, anisotropic magnification and higher-order aberrations. Beam-induced particle motions were corrected using Bayesian polishing module in RELION. The resulting ‘shiny’ particles were refined with mask2, resulting in a final 3.2 Å resolution reconstruction of the H/ACA RNP module (Extended Data Fig. 2a, b, d).

For the catalytic core, the combined 2,614,217 sub-particles (pixel size of 2.2 Å) were refined to 4.4 Å resolution using mask1 (Extended Data Fig. 1d), and then classified into six classes (RELION options: --sigma\_ang 20 --healpix\_order 3). A total of 1,386,006 sub-particles from the two best classes were combined, extracted in boxes of 256 pixels square (pixel size of 1.1 Å), and refined to 3.4 Å resolution. Notably, the cryo-EM densities of the TEN domain of TERT and P2a.1–P2a of TER were weaker than the rest in the resulting map, suggesting that they are positionally flexible and/or adopt more than one conformation. The sub-particles were therefore subjected to another round of 3D classification with a finer angular sampling (RELION options: --sigma\_ang 6 --healpix\_order 4) and a tighter soft mask (mask3) to exclude the flexible regions described above. 291,504 sub-particles from the best class were selected for 3D refinement, CTF refinement and Bayesian polishing. The resulting ‘shiny’ sub-particles were refined to 3.3 Å resolution with mask1 for the entire catalytic core and to 3.2 Å resolution with mask3 for the rigid region that excludes TEN and P2a.1–P2a (Extended Data Fig. 2a, c, e). Several alignment-free 3D classifications

using different masks (Extended Data Fig. 3) were subsequently performed to investigate the compositional and conformational heterogeneities within the catalytic core, as described below.

To separate sub-particles bound with H2A–H2B heterodimer, a soft mask covering the junction region of P2b, P6.1 and P6 was used in the 3D classification (Extended Data Fig. 3a). The input 291,504 sub-particles were classified into 10 classes with an optimized regularization parameter (RELION option: --tau2\_fudge 8). A subset of 65,345 sub-particles (21.9% of the inputs) in one class with clear features for H2A–H2B were selected and refined to 3.7 Å resolution using mask1 (Extended Data Fig. 3a).

To separate sub-particles bound with TPP1, the inputs were classified into 4 classes with a spherical soft mask covering the space above the TEN-TRAP (Extended Data Fig. 3a). A subset of 118,845 sub-particles in one class with the best density for TPP1 were selected and refined to 3.5 Å resolution (Extended Data Fig. 3a, c, d). Meanwhile, 83,992 sub-particles from another class without TPP1 were refined to 3.7 Å resolution. These two subsets of catalytic core sub-particles were further classified into six classes, respectively, with a soft mask encasing the P2a.1–P2a of TER (Extended Data Fig. 3a). The resulting classes with interpretable features (that is, major and minor grooves of the RNA duplex) were refined individually.

Resolutions of the cryo-EM maps were estimated on the basis of the gold standard<sup>59</sup> Fourier shell correlation (FSC) = 0.143 criterion. The cryo-EM maps were corrected for the modulation transfer function of the detector, sharpened with a negative B-factor and low-pass filtered to the stated resolution using the `relion_postprocess` program in RELION. Local resolution evaluations were determined by RELION with two independently refined half-maps. Data collection and processing statistics are given in Extended Data Table 1.

## Reference models

The initial model for 3D reconstruction was retrieved from the EMDB with accession code EMD-7521. Other structures used in this study were retrieved from the PDB with accession codes 2I46 (TPP1 OB), 7LMA (*Tetrahymena* telomerase), 7BG9 (human telomerase catalytic core), 7BGB (human telomerase H/ACA RNP), and 5CQG (*Tribolium* TERT-like protein with inhibitor).

## Model building and refinement

The atomic models of human telomerase catalytic core were built and refined manually in COOT<sup>60</sup>. The 3.3 Å resolution cryo-EM map of catalytic core (Extended Data Fig. 1d) was used for the modelling of TERT RBD-RT-CTE ring and associated TER CR4/5 and t/PK except for the P2a.1 and P2a. The 3.5 Å resolution map of catalytic core bound with TPP1 was used for the modelling of TERT TEN-TRAP, TER P2a and P2a.1, as well as TPP1. The 3.7 Å resolution map of catalytic core bound with H2A–H2B was used for the modelling of H2A and H2B.

Initially, our model of human telomerase catalytic core<sup>36</sup> based on a reported 8 Å resolution cryo-EM map<sup>9</sup> was fitted into our catalytic core maps with UCSF Chimera<sup>61</sup> as a beginning

for model building. The high-resolution features of our maps provided more accurate main chain and most side chain conformations (Extended Data Fig. 2g–j), which aided the side chain adjustment and main chain re-tracing when necessary to get the best fit between the density and the model. Visible densities of amino acid residues with bulky side chains, such as Phe, Tyr and Trp were used as guidance for sequence assignment. No density was observed for the linker between TEN and RBD (residues 179–321), RBD residues 417–443, and RT residues 641–650, so these residues were not modelled. The crystal structure of TPP1 OB<sup>33</sup> (PDB: 2146) was fitted into the telomerase-TPP1 sub-class maps and manually refined in a similar way (Extended Data Fig. 3f). The H2A–H2B heterodimer (Extended Data Fig. 3b) was adapted from the structure of human telomerase holoenzyme<sup>8</sup> (PDB: 7BG9) published during the preparation of this manuscript. Comparison of our model with the published structure revealed some differences in residue register (about 16% overall for visible regions of TERT), especially within the TEN (37%). Our telomerase–TPP1 structure exhibits less positional dynamics and therefore higher resolution for TEN–TRAP, while the higher resolution of the catalytic core in the combined maps provided better density for backbone and sidechain positioning for the TERT ring and associated TER.

TER regions contacting TERT, especially the t/PK domain and P6.1 stem, were de novo built from the well-defined nucleotide densities (Extended Data Fig. 2i). Other parts of TER were adjusted from our previous model<sup>36</sup> for their base conformations and backbones to fit into the density map. The different conformations of the most flexible region of TER (P2a.1–P2a) were achieved from different classes of local refined maps in a similar way (Extended Data Fig. 3a). The substrate DNA was traced from the 3' end inside the catalytic cavity with seven nucleotides modelled including the G18–A15 which form a four base pair duplex with the template. Nucleotides T1–T11 were invisible in the cryo-EM map.

For the modelling of H/ACA RNP, the previously published model<sup>8</sup> (PDB: 7BGB) was used as the starting point. The model was rigid-body fitted into our 3.2 Å resolution cryo-EM map of H/ACA RNP and adjusted manually in COOT.

Catalytic core and H/ACA RNP models were refined individually using Phenix<sup>62</sup> in real space with secondary structure, Ramachandran and rotamer restraints. All structures were validated using the Molprobity scores and statistics of the Ramachandran plots. Refinement statistics of the models are summarized in Extended Data Table 1. Model vs map FSC validation is shown in Extended Data Figs. 2, 3. All figures presenting the model were prepared using UCSF Chimera<sup>61</sup>, ChimeraX<sup>63</sup> and PyMOL<sup>64</sup>.

### **In vitro reconstitution of telomerase and direct telomerase activity assays**

Human TER for telomerase activity assays was prepared as described for *Tetrahymena* TER<sup>10</sup>, with minor changes. In brief, TER was transcribed in vitro with homemade T7 RNA polymerase (P226L mutant)<sup>65</sup> using 1 μM linearized pUC19-T7-TER template in a reaction containing 40 mM Tris·HCl pH 8.0, 5 mM of each dNTP, 40 mM MgCl<sub>2</sub>, 1 mM spermidine, 2.5 mM DTT, and 0.01% Triton X-100. The reaction was performed at 37 °C for 4 h. Product was purified with a 10% denaturing polyacrylamide gel followed by electroelution. Human telomerase catalytic core (TERT and TER) was reconstituted in RRL as described<sup>44,66</sup> with minor changes. Wild type (WT) and mutant human TERTs were

expressed in RRL using the TNT quick coupled transcription/translation system (Promega) for 2 h before incubation with 1  $\mu$ M TER with and without 10  $\mu$ M TPP1 (or 0.25  $\mu$ M TPP1–POT1 as noted) for another 30 min. The concentrations of TPP1 and TPP1–POT1 used in the assays were optimized under linear conditions. Assays were performed in a 20  $\mu$ l reaction volume containing 25 mM Tris-HCl pH 8.3, 2 mM MgCl<sub>2</sub>, 1 mM DTT, 500  $\mu$ M dTTP, 500  $\mu$ M dATP, 12.5  $\mu$ M dGTP, and 3–5  $\mu$ Ci  $\alpha$ -<sup>32</sup>P-dGTP (Perkin-Elmer), 500 nM DNA primer d(TTAGGG)<sub>3</sub> and 10  $\mu$ l of telomerase reconstituted from RRL. After 1 h incubation at 37 °C, reactions were stopped by adding 80  $\mu$ l quench buffer (10 mM Tris-HCl, pH 8.0 and 10 mM EDTA) supplied with a 15-nt <sup>32</sup>P end-labelled primer as a recovery control. Products were extracted with phenol-chloroform, ethanol precipitated, and resolved on a 10% denaturing polyacrylamide gel. The gel was vacuum dried and exposed to a phosphor imaging screen for 20 h. The final image was obtained by scanning the screen on a ImageFX system (Bio-Rad) and then quantified with QuantityOne (Bio-rad). Telomerase activity and RAP were determined as previously described<sup>10</sup>. The integrated intensity of each lane was used for calculation of telomerase activity and intensities of bands representing the extension of one telomere repeat processed using the fraction left behind (FLB) method<sup>67,68</sup> for calculation of telomerase RAP (Supplementary Fig. 1). WT TERT without TPP1 was set as 1 for both activity and RAP, while WT with TPP1 along with all other mutants were compared to it for relative activity and RAP. Activity stimulation was calculated as relative activity with TPP1/relative activity without TPP1. Activity stimulation loss was calculated as (activity stimulation of WT TERT – activity stimulation of mutant TERT)/[(activity stimulation of WT TERT) – 1].

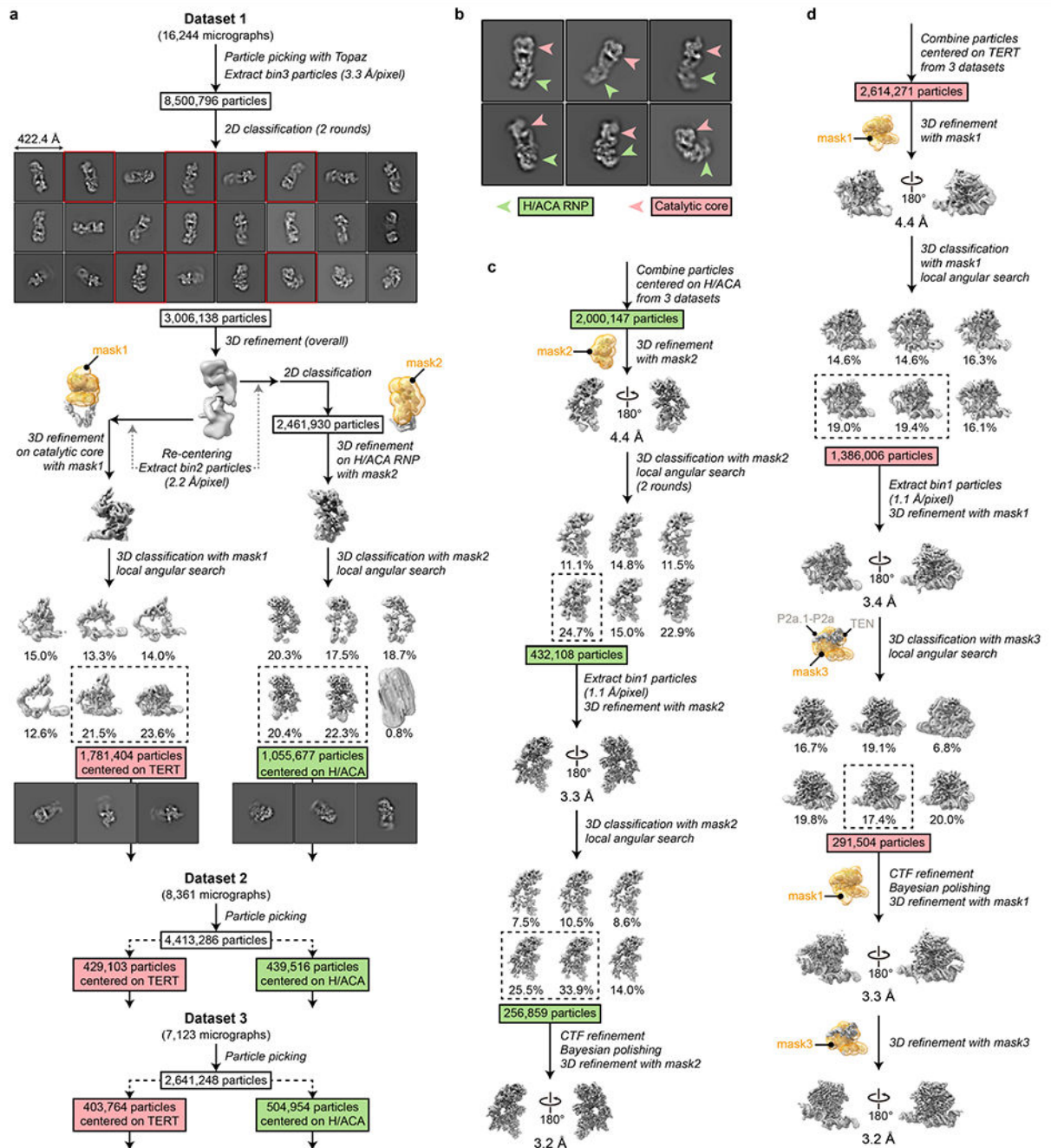
### Reporting summary

Further information on research design is available in the Nature Research Reporting Summary linked to this paper.

### Data availability

Cryo-EM density maps have been deposited in the Electron Microscopy Data Bank under accession numbers EMD-26085 (H/ACA RNP), EMD-26086 (catalytic core combined), EMD-26087 (catalytic core with TPP1), EMD-26096 (catalytic core without TPP1), EMD-26088 (catalytic core with H2A–H2B), EMD-26090 (catalytic core with TPP1, P2a state 1-1), EMD-26091 (catalytic core with TPP1, P2a state 1-2), EMD-26092 (catalytic core with TPP1, P2a state 1-3), EMD-26093 (catalytic core with TPP1, P2a state 2), EMD-26094 (catalytic core without TPP1, P2a state 1) and EMD-26095 (catalytic core without TPP1, P2a state 2). The atomic models have been deposited in the Protein Data Bank under accession codes 7TRC (H/ACA RNP), 7TRD (catalytic core combined), 7TRE (catalytic core with TPP1) and 7TRF (catalytic core with H2A–H2B). Uncropped version of all the gels are included as Supplementary Fig. 1. Any other relevant data are available from the corresponding authors upon reasonable request.

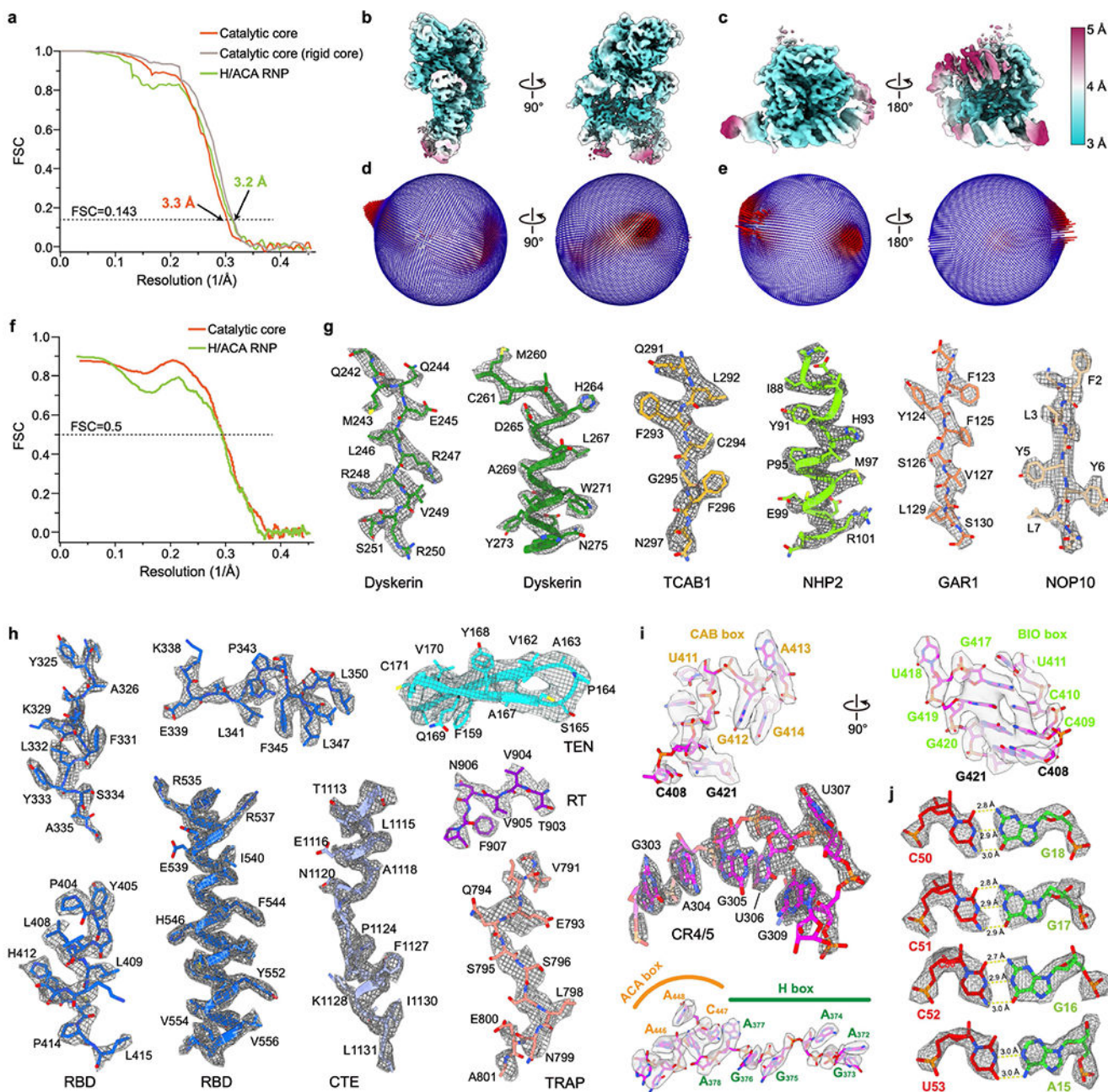
## Extended Data



**Extended Data Fig. 1 | Cryo-EM data processing workflow of human telomerase (detailed in Methods).**

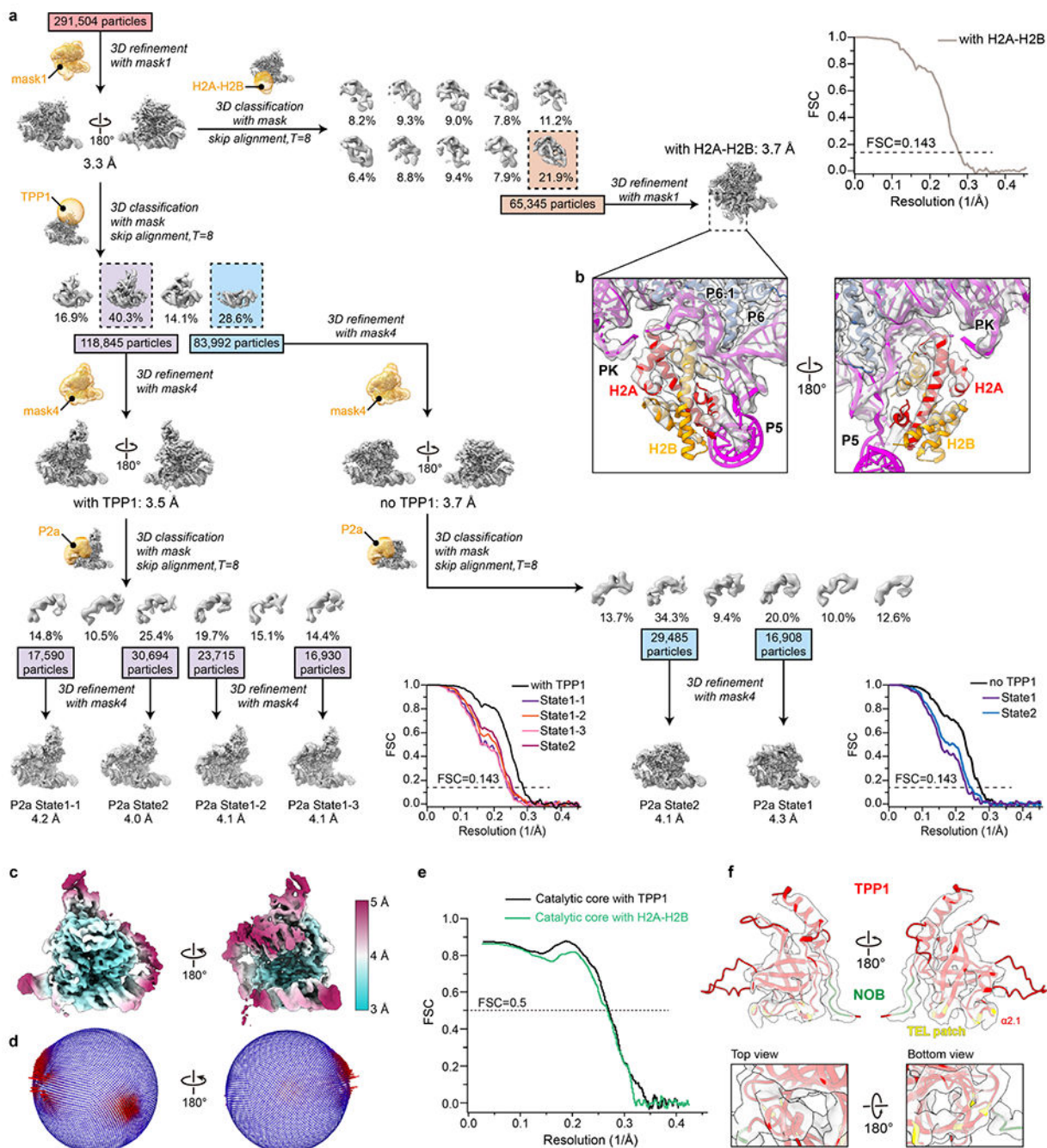
**a**, Initial screening of sub-particles centered on the catalytic core and H/ACA RNP. **b**, Representative 2D class images of human telomerase as indicated in the red boxes in **a**. **c**, **d**, 3D classification and reconstruction processes for the H/ACA RNP (**c**) and catalytic core (**d**). Soft masks used in the data processing are shown in orange.





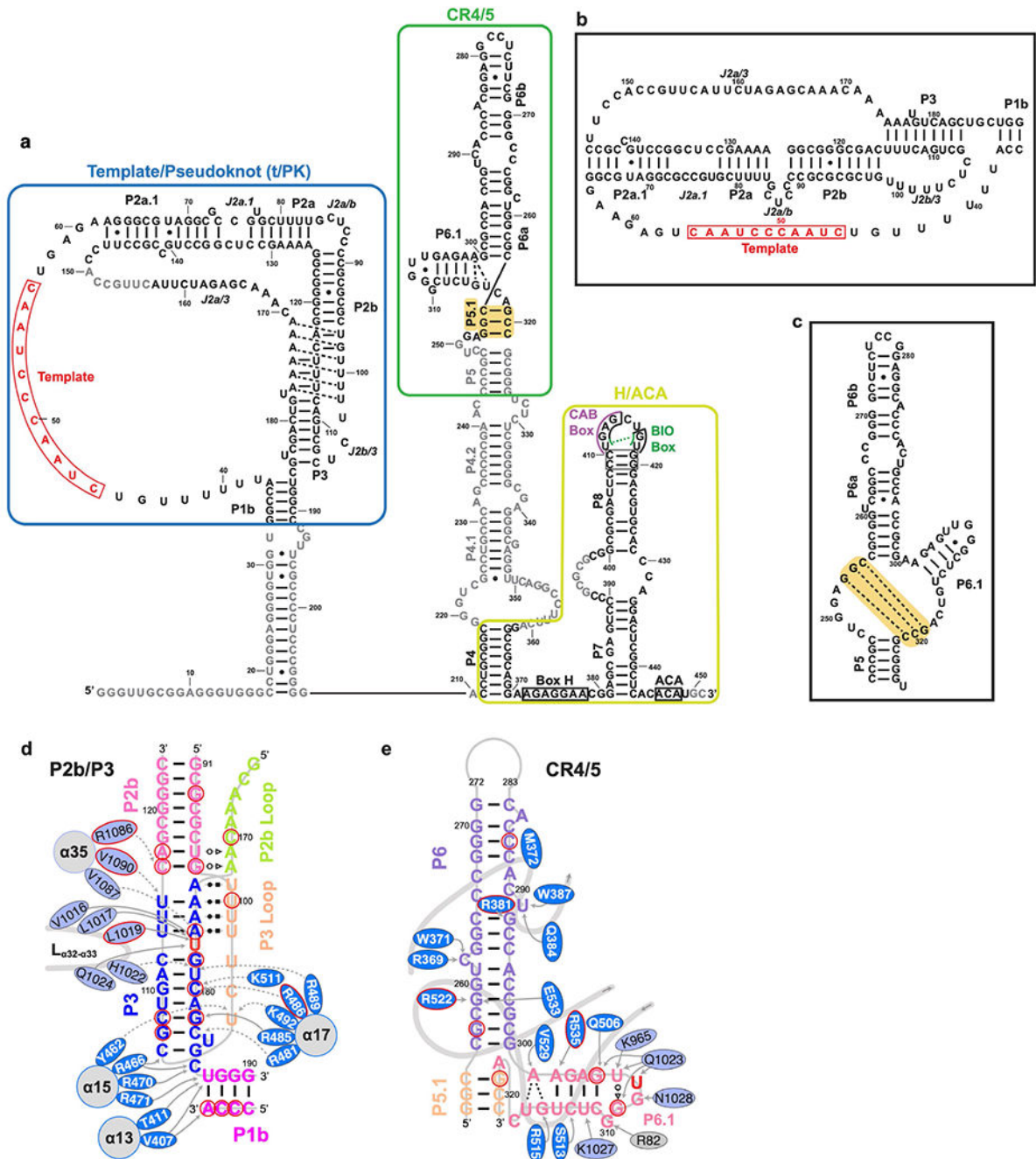
**Extended Data Fig. 2 | Evaluation of cryo-EM reconstructions.**

**a**, Plot of the Fourier shell correlation (FSC) as a function of the spatial frequency of the final reconstructions. **b, c**, Local resolution evaluation of the H/ACA RNP (**b**) and catalytic core (**c**) reconstructions. **d, e**, Euler angle distribution of sub-particles used for the H/ACA RNP (**d**) and catalytic core (**e**) reconstructions. **f**, FSC coefficients as a function of spatial frequency between model and corresponding cryo-EM density maps. **g–j**, Representative cryo-EM densities encasing the corresponding atomic models of H/ACA proteins (**g**), TERT (**h**), TER (**i**) and the four base pairs in the template–DNA duplex (**j**).



### Extended Data Fig. 3 | Analysis of compositional and conformational heterogeneity of the catalytic core.

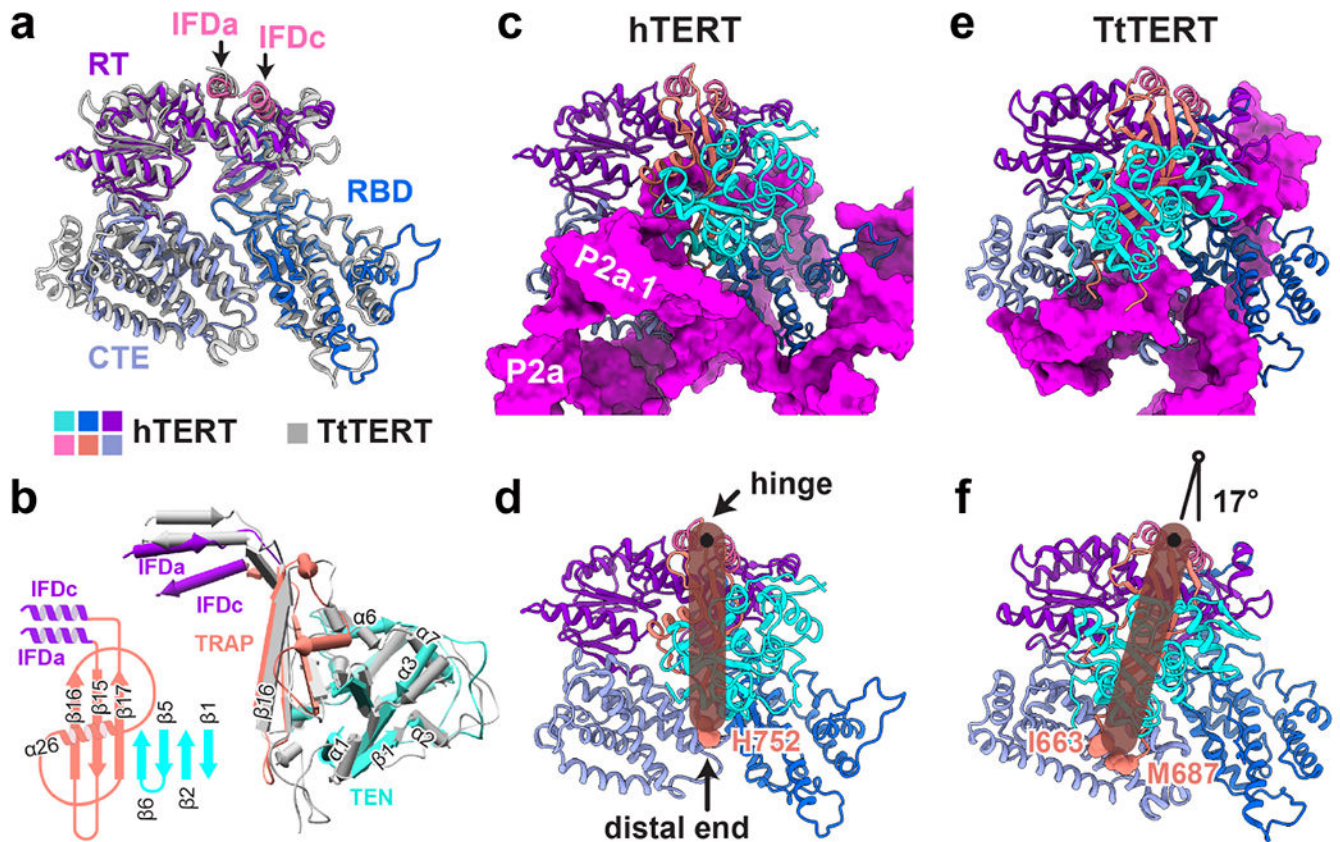
**a**, Cryo-EM 3D classification and reconstruction workflow with different masks. Insets show the FSC curves of final reconstructions. **b**, Superposition of H2A-H2B model and cryo-EM density. **c**, Local resolution evaluation of the 3.5 Å resolution reconstruction of catalytic core bond with TPP1. **d**, Euler angle distribution of sub-particles used for the reconstruction of catalytic core bound with TPP1. **e**, Model VS map FSC curve of the reconstruction with TPP1 or H2A-H2B. **f**, Superposition of TPP1 model and cryo-EM density.



#### Extended Data Fig. 4 | Secondary structure of human TER and interactions with TERT.

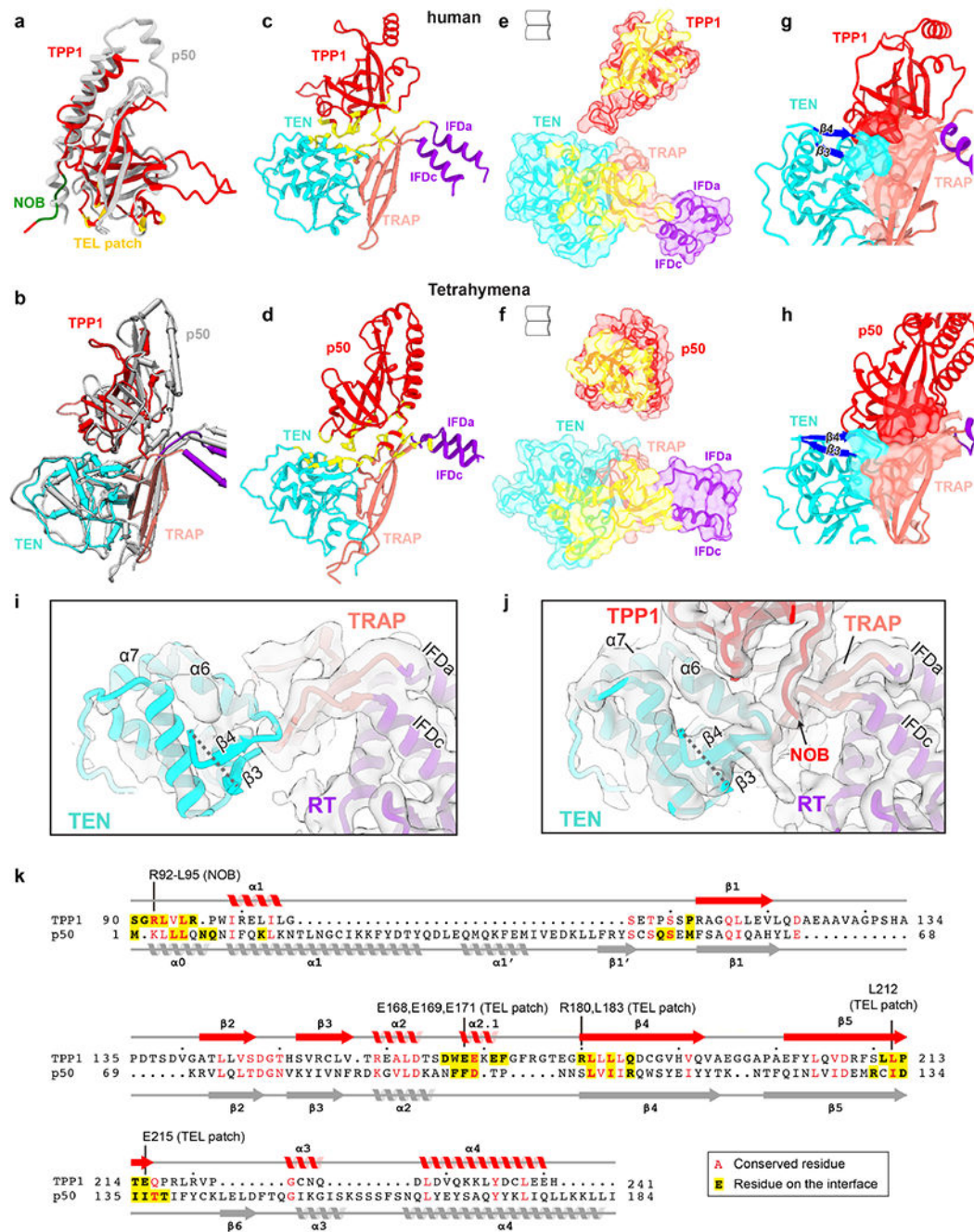
**a**, Secondary structure observed in our telomerase structure (black), with unmodeled regions (gray) based on the telomerase database<sup>13</sup>. **b, c**, The previously predicted secondary structures of t/PK (**b**) and CR4/5 (**c**) based on phylogenetic analysis<sup>13</sup>. The dashed lines connecting residues in **c** indicate interactions forming the 3 bp P5.1 (highlighted) found in our structure, added for clarity. **d, e**, Structure-based schematics of TER P2b/P3-P1b (**d**) and CR4/5 (**e**) and interactions with TERT. Solid and dashed arrows from TERT residues to TER nucleotides indicate interactions with base and backbone, respectively. Base pair types are

indicated using Leontis-Westhof symbols<sup>69</sup>. Red circles/ovals indicate residue positions with disease related mutations.



**Extended Data Fig. 5 | Comparison of human and *Tetrahymena* TERT structure.**

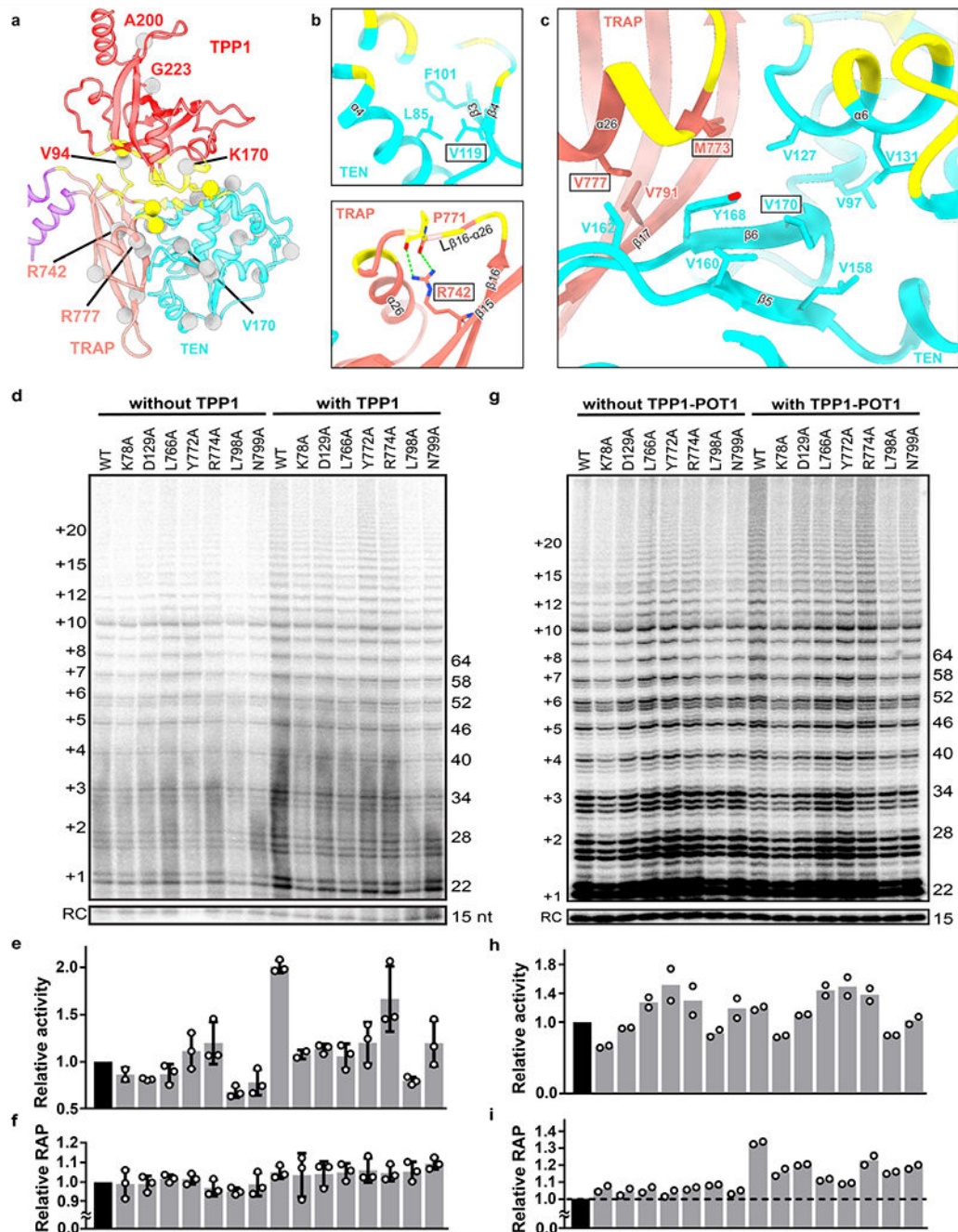
**a**, Superposition of RBD-RT-CTE ring (TERT ring) structures of hTERT and *Tetrahymena* TERT (TtTERT, PDB: 7LMA). **b**, Overlay of TEN-IFD-TRAP structures in human (colored) and *Tetrahymena* (gray) telomerase. Inset shows the secondary structure of human TRAP and the extended  $\beta$  sheet between TEN and TRAP. **c-f**, Comparison of human and *Tetrahymena* TEN-TRAP position relative to the TERT ring. Ribbon diagrams of the TERT rings from hTERT (**c-d**) and TtTERT (**e-f**) are shown in the same orientations as in **A**. Human and *Tetrahymena* TER are shown as magenta surfaces in **c** and **e**, respectively. Transparent sticks in **d** and **f** indicate the orientations of TRAP domains in hTERT and TtTERT, respectively. Residues located at the distal end of TRAP domains are shown as spheres. The hinge of the “rotation movement” is at the proximal end of TRAP that is connected to the IFDa and IFDc helices.



**Extended Data Fig. 6 | Comparison of human TEN-TRAP-TPP1 and *Tetrahymena* TEN-TRAP-p50.**

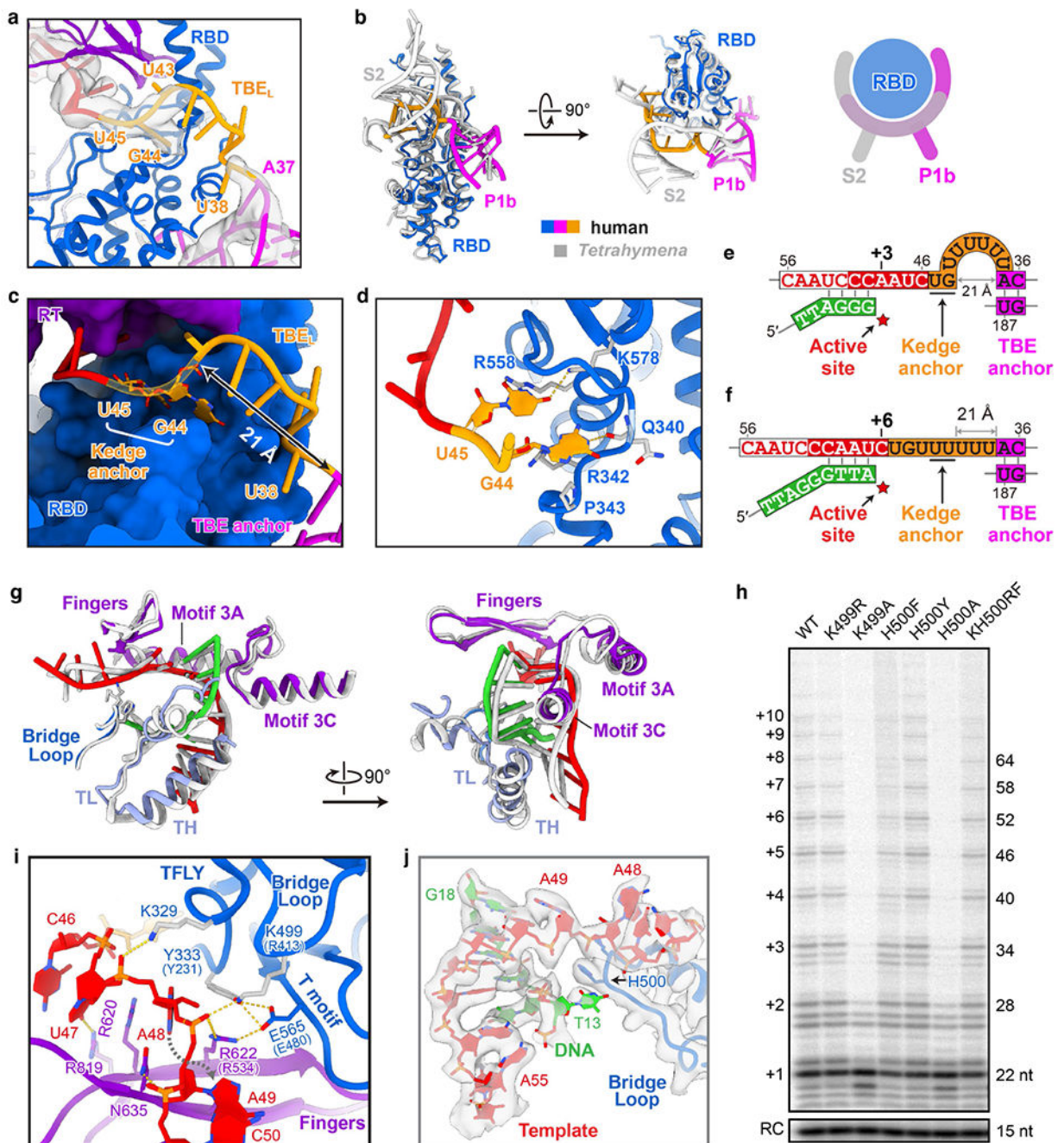
**a**, Comparison of telomerase-bound TPP1 and p50 (gray) (PDB: 7LMA). **b**, Superposition of human TEN-TRAP-TPP1 (colored as in Fig. 2) and *Tetrahymena* TEN-TRAP-p50 (gray). **c**, **d**, Overall comparison of human TEN-TRAP-TPP1 (**c**) and *Tetrahymena* TEN-TRAP-p50 (**d**) with the residues locate on the interfaces colored yellow. **e**, **f**, Open book views of the continuous interfaces between human TPP1-TERT (**e**) and *Tetrahymena* p50-TERT (**f**) shown as surface, respectively. **g**, **h**, A three-way junction shared by human

TEN-TRAP-TPP1 (g) and *Tetrahymena* TEN-TRAP-p50<sup>10</sup> (h). i, j. Density maps of TEN and TRAP without (i) and with (j) TPP1, highlighting the changes in density upon TPP1 binding. Density thresholds were adjusted for comparable density in the RT region. k. Structure-based sequence alignment (rendered with ESPrict3<sup>70</sup>) for human TPP1 (red) and *Tetrahymena* p50 (grey). Residues located on the interface are highlighted in yellow. Residues comprising TPP1 NOB and TEL patch are indicated on the top of alignment.



Extended Data Fig. 7 | Disease related mutations and telomerase activity assays.

**a-e** Locations of disease related mutations. **a**, Back view of TEN–TRAP–TPP1 interface as in Extended Data Fig. 6. Residue positions with disease mutations in TEN, TRAP, and TPP1 (Supplementary Table 1<sup>13,71,72</sup>) are shown as yellow (interface) and gray (non-interface) spheres. **b, c**, Zoomed-in views of intra-(**b**) and inter-(**c**) domain interactions from residues related to disease mutations that may disrupt TPP1–TERT interaction indirectly. **d–i** Telomerase activity assays for TERT variants with residue substitutions at the interface with TPP1. **d**, Telomerase activity assays corresponding to Fig. 2g. Gel is a representative from 3 independent experiments. The number of telomeric repeats added to primer are indicated at left, and number of nucleotides are indicated at right. RC, recovery control. **e, f**, Relative activity (**e**) and RAP (**f**) normalized to TERT WT without TPP1. Plotted values are mean  $\pm$  s.d. from  $n = 3$  biologically independent experiments. **g**, Telomerase activity assays without and with TPP1–POT1. Gel is a representative from 2 independent experiments. **h, i**, Relative activity (**h**) and RAP (**i**) normalized to TERT WT without TPP1–POT1. Open circles are values from  $n = 2$  biologically independent experiments.

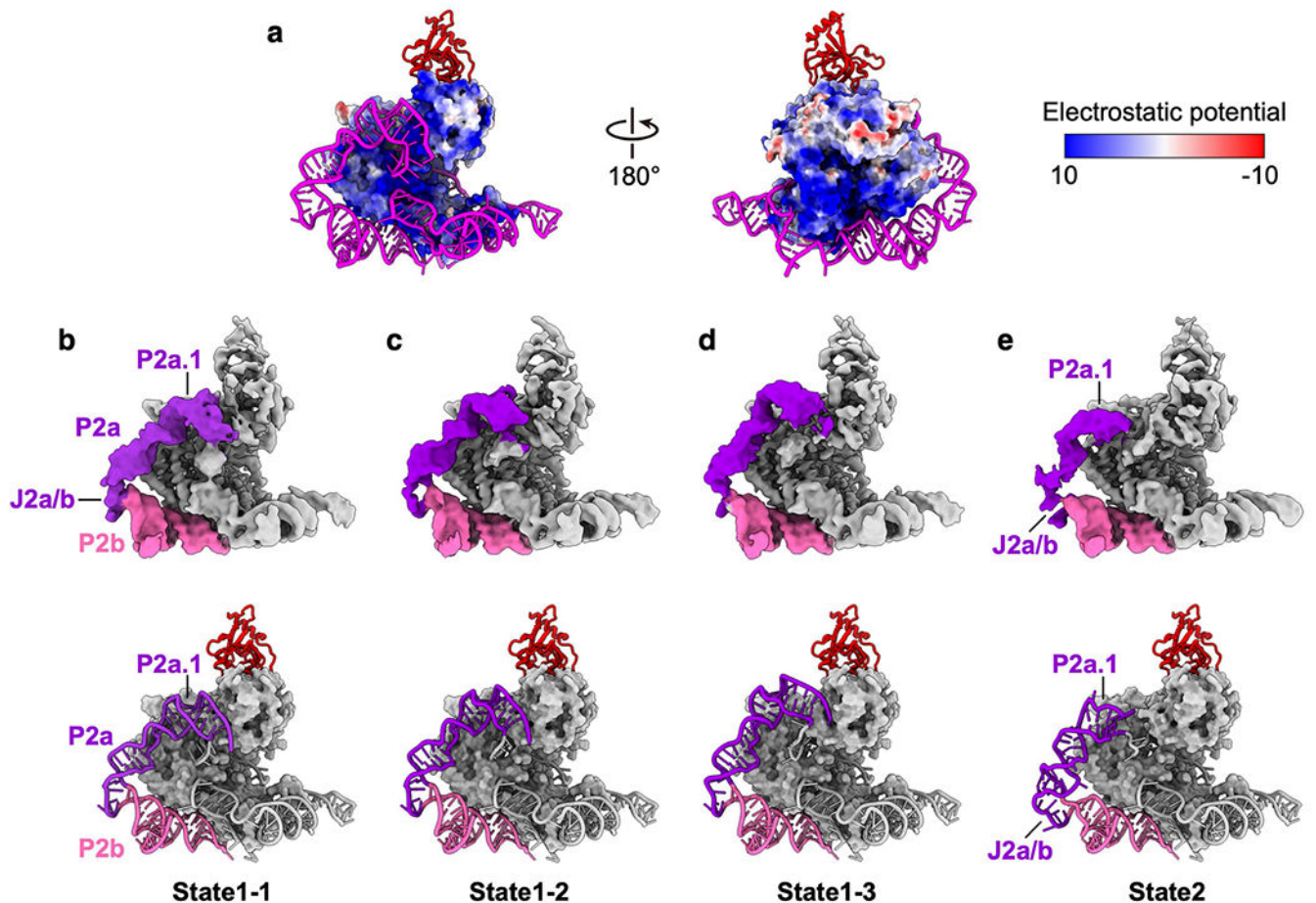


### Extended Data Fig. 8 | Structural details of the TBE, bridge loop and related activity assays.

**a**, Structure of the single-stranded RNA nucleotides at the 5' side of the template in complex with TERT RBD (colored as in Fig. 3a). Cryo-EM densities of TER nucleotides are overlaid on the structure as transparent surfaces. Only the first two template-adjacent nucleotides (G<sub>44</sub>U<sub>45</sub>) and P1b have strong densities in the cryo-EM map, whereas the intervening nucleotides show only weak density indicative of positional dynamics. **b**, Comparison of the TBE-TBE<sub>L</sub>-RBD structures from human (color) and *Tetrahymena* (gray, PDB: 7LMA) telomerase. **c**, G<sub>44</sub> and U<sub>45</sub> in the kedge anchor pocket of TERT RBD.



The linear distance from the phosphate group of U<sub>38</sub> to the phosphate group of G<sub>44</sub> is about 21 Å. **d**, Detailed interactions between G<sub>44</sub>-U<sub>45</sub> and TERT RBD. Intermolecular hydrogen bonds are shown as dashed yellow lines. **e**, Schematic of TER TBE-TBE<sub>L</sub>-template conformation when the template is at the +3 position as in our structure. **f**, Predicted TBE-TBE<sub>L</sub>-template conformation when the template is at the +6 position. TBE<sub>L</sub> nucleotides U<sub>38–40</sub> would be fully stretched to span the distance (21 Å) from the TBE anchor to the kedge anchor. **g**, Comparison of the template–DNA duplex and surrounding structural elements of human (color) and *Tetrahymena* (gray, PDB: 7LMA) telomerase. Residues located on the tip of the bridge loop are shown as sticks. **h**, Telomerase activity assays with substitutions of hTERT K499 and/or H500. Gel is a representative from 2 independent experiments. **i**, Detailed interactions surrounding the entrance of template nucleotides. Side chains of key residues are indicated with corresponding *Tetrahymena* TERT residues in parentheses. **j**, Ribbon diagram of the template–DNA duplex and the bridge loop superimposed with cryo-EM densities (transparent surface).



**Extended Data Fig. 9 | Electrostatic interactions and conformational dynamics of the pseudoknot on TERT.**

**a**, Electrostatic surface of TERT shown in two different views. **b–e**, Cryo-EM density (upper) and model (lower) of telomerase catalytic core with P2 stem in State1-1 (**b**), State1-2 (**c**), State1-3 (**d**) and State2 (**e**). Through State1-1 to 1-3 (**b–d**), P2a.1 conducts

an upward movement. From State1-1 (b) to State2 (e), P2a.1 and P2a move away from TEN domain of TERT with the J2a/b linker bulged out. The conformation of P2b and its location on TERT remain the same in the four structures.

### Extended Data Table 1 |

Cryo-EM data collection, refinement and validation statistics for human telomerase maps and models

	H/ACA RNP (EMD-26085) (PDB 7TRC)	Catalytic core combined (EMD-26086) (PDB 7TRD)	Catalytic core with TPP1 (EMD-26087) (PDB 7TRE)	Catalytic core without TPP1 (EMD-26096)	Catalytic core with H2A-H2B (EMD-26088) (PDB 7TRF)
<b>Data collection and processing</b>					
Magnification	81,000	81,000	81,000	81,000	81,000
Voltage (kV)	300	300	300	300	300
Electron exposure (e <sup>-</sup> /Å <sup>2</sup> )	55	55	55	55	55
Defocus range (μm)	-0.8 – -4.0	-0.8 – -4.0	-0.8 – -4.0	-0.8 – -4.0	-0.8 – -4.0
Pixel size (Å)	1.1	1.1	1.1	1.1	1.1
Symmetry imposed	C1	C1	C1	C1	C1
Initial particle images (no.)	15,555,330	15,555,330	15,555,330	15,555,330	15,555,330
Final particle images (no.)	256,859	291,504	118,845	83,992	65,345
Map resolution (Å)	3.2	3.3	3.5	3.7	3.7
FSC threshold	0.143	0.143	0.143	0.143	0.143
Map resolution range (Å)	3.2–6.0	3.3–6.0	3.5–7.0	3.7–7.0	3.7–7.0
<b>Refinement</b>					
Initial model used (PDB code)	7BGB		2I46		7BG9
Model resolution (Å)	3.4	3.4	3.6		3.7
FSC threshold	0.5	0.5	0.5		0.5
Map sharpening <i>B</i> factor (Å <sup>2</sup> )	-106.0	-100	-100		-50
<b>Model composition</b>					
Non-hydrogen atoms	14,747	12,240	13,411		13,581
Protein residues	1,599	934	1,086		1,106
RNA/DNA Nucleotides	92	224	224		224
Ligands	0	0	0		0
<b>R.m.s. deviations</b>					
Bond lengths (Å)	0.003	0.004	0.006		0.005
Bond angles (°)	0.645	0.622	0.757		0.734
<b>Validation</b>					
MolProbity score	1.74	1.71	1.82		1.73
Clashscore	8.21	7.37	7.28		7.36

	H/ACA RNP (EMD-26085) (PDB 7TRC)	Catalytic core combined (EMD-26086) (PDB 7TRD)	Catalytic core with TPP1 (EMD-26087) (PDB 7TRE)	Catalytic core without TPP1 (EMD-26096)	Catalytic core with H2A-H2B (EMD-26088) (PDB 7TRF)
Poor rotamers (%)	0.21	0.12	0.11		0.10
Ramachandran plot					
Favored (%)	95.80	95.67	93.58		95.33
Allowed (%)	4.20	4.33	6.42		4.58
Disallowed (%)	0.00	0.00	0.00		0.09

	Catalytic core with TPP1, P2a State1-1 (EMD-26090)	Catalytic core with TPP1, P2a State 1-2 (EMD-26091)	Catalytic core with TPP1, P2a State 1-3 (EMD-26092)	Catalytic core with TPP1, P2a State2 (EMD-26093)	Catalytic core without TPP1, P2a State 1 (EMD-26094)	Catalytic core without TPP1, P2a State2 (EMD-26095)
<b>Data collection and processing</b>						
Magnification	81,000	81,000	81,000	81,000	81,000	81,000
Voltage (kV)	300	300	300	300	300	300
Electron exposure (e <sup>-</sup> /Å <sup>2</sup> )	55	55	55	55	55	55
Defocus range (μm)	-0.8 – -4.0	-0.8 – -4.0	-0.8 – -4.0	-0.8 – -4.0	-0.8 – -4.0	-0.8 – -4.0
Pixel size (Å)	1.1	1.1	1.1	1.1	1.1	1.1
Symmetry imposed	C1	C1	C1	C1	C1	C1
Initial particle images (no.)	15,555,330	15,555,330	15,555,330	15,555,330	15,555,330	15,555,330
Final particle images (no.)	17,590	23,715	16,930	30,694	16,908	29,485
Map resolution (Å)	4.2	4.1	4.1	4.0	4.3	4.1
FSC threshold	0.143	0.143	0.143	0.143	0.143	0.143
Map resolution range (Å)	4.2–8.0	4.1–8.0	4.1–8.0	4.0–8.0	4.3–8.0	4.1–8.0

## Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

## Acknowledgements

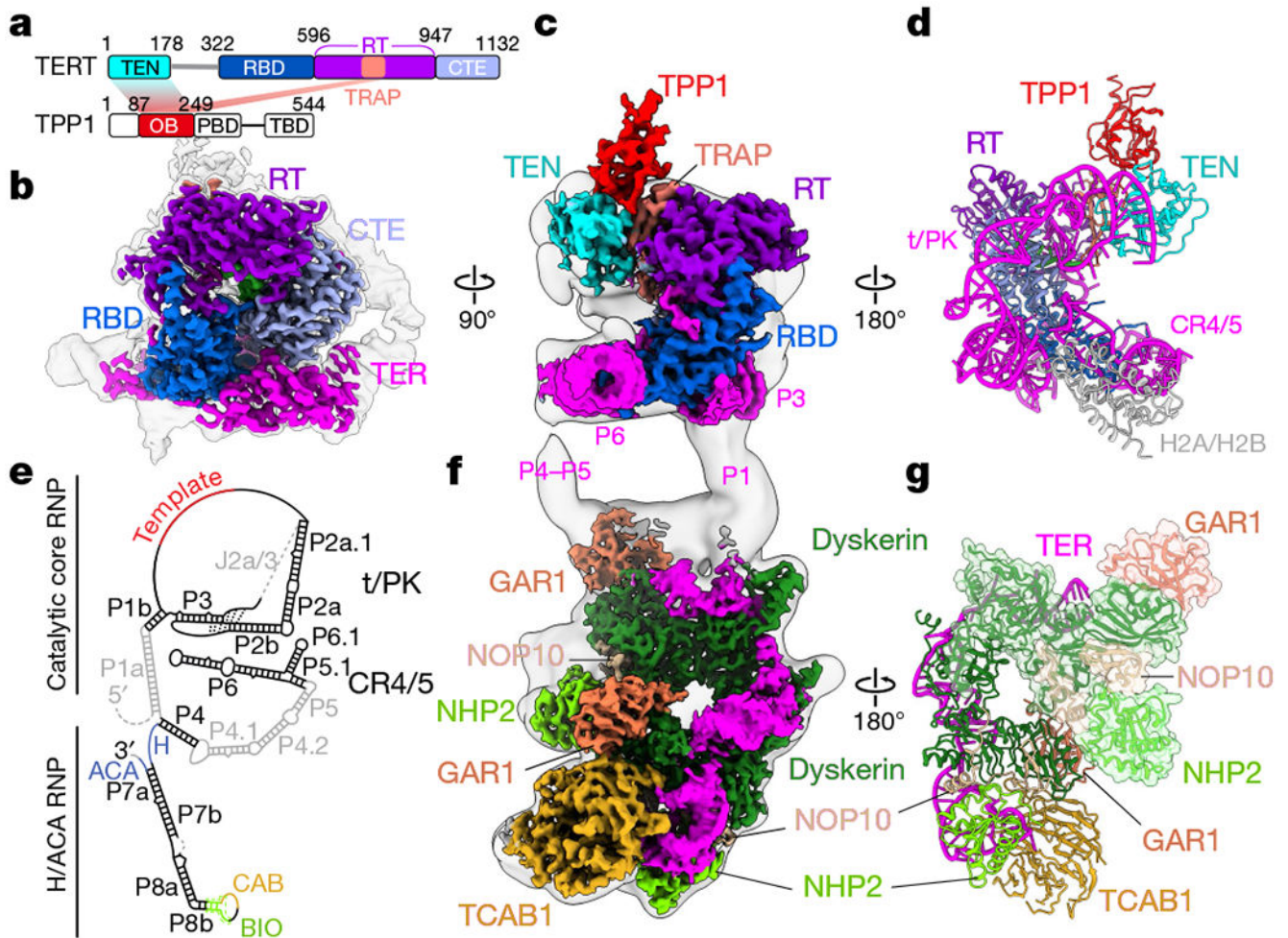
This work was supported by grants from NIH R35GM131901 and NSF MCB2016540 to J.F. and NIH R01GM071940 to Z.H.Z. We acknowledge use of instruments at the Electron Imaging Center for Nanomachines supported by UCLA and instrumentation grants from NIH (1S10RR23057, 1S10OD018111) and NSF (DBI-1338135 and DMR-1548924).

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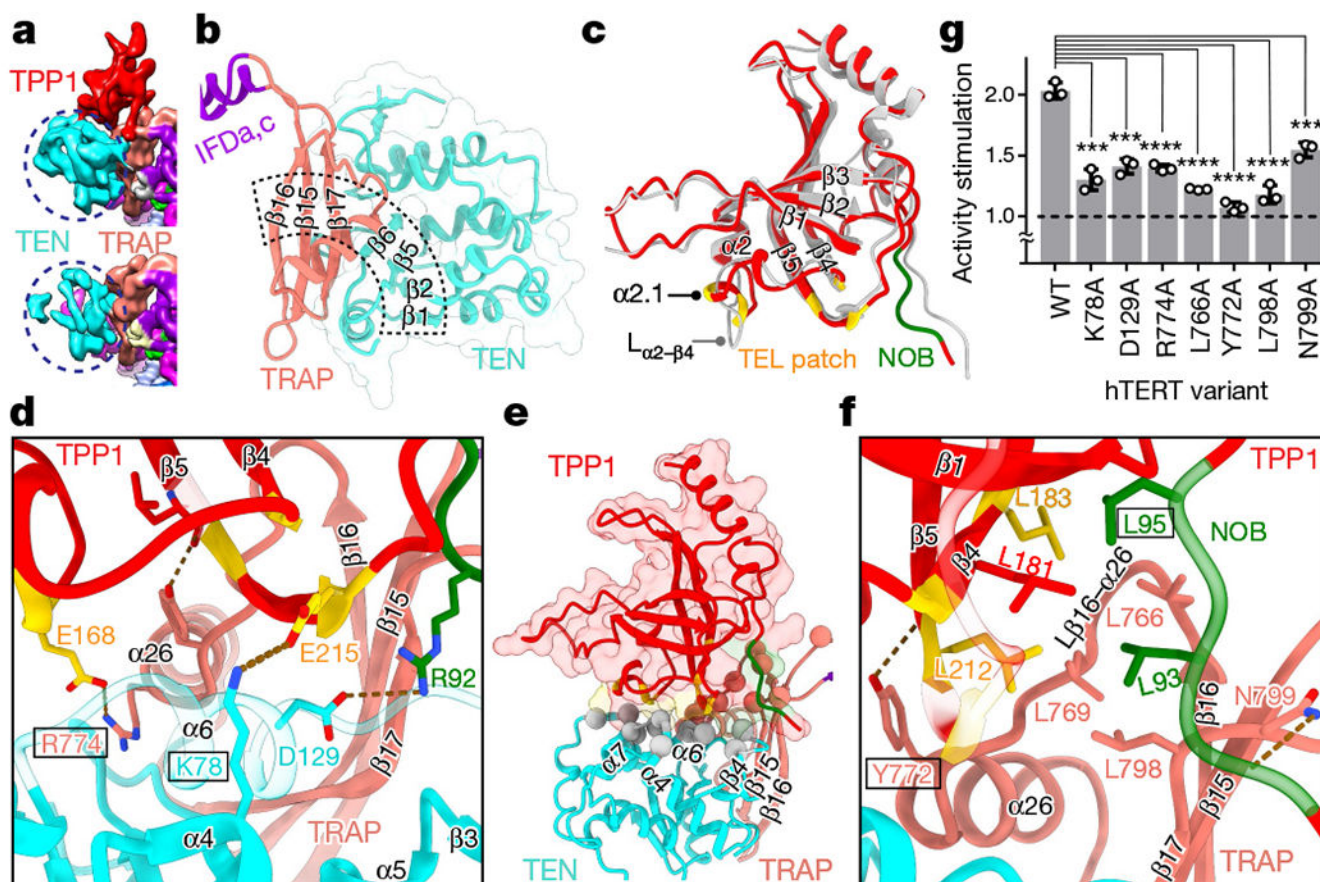
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**Fig. 1 | Structure of human telomerase with TPP1.**

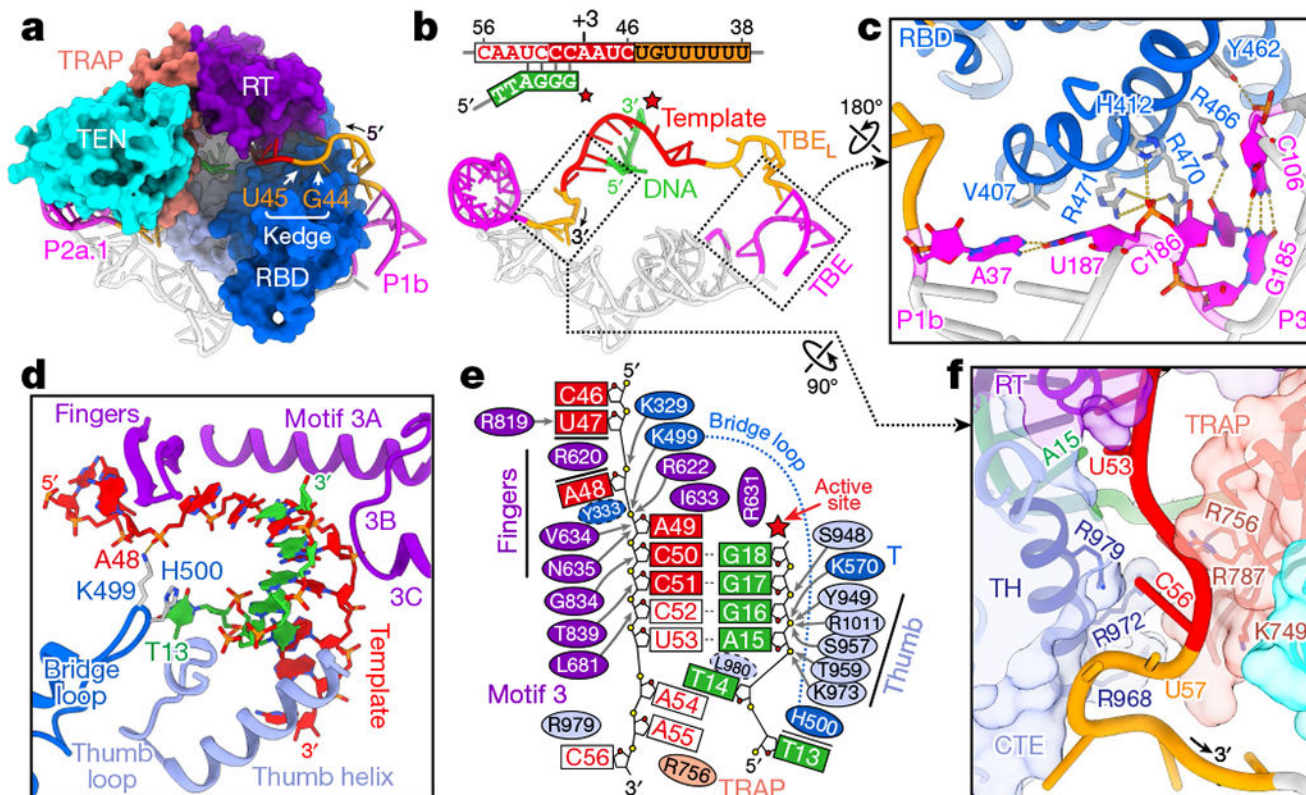
**a**, Schematic of TERT and TPP1 domains and interactions. PBD, POT1-binding domain; TBD, TIN2-binding domain. **b**, Cryo-EM density map of catalytic core at 3.3 Å resolution shown at two different thresholds. The high-threshold map showing high-resolution features is coloured as in **a** for TERT and magenta for TER, and the low-threshold map showing the continuous density is overlaid as transparent surface. **c**, The 3.5 Å-resolution map of the catalytic core bound with TPP1. **d**, Cartoon model of the catalytic core. **e**, Schematic of TER secondary structure determined in the model. Regions invisible in the cryo-EM map are shown in grey. **f**, **g**, Cryo-EM density map (**f**) and cartoon model (**g**) of H/ACA RNP. Cryo-EM map of the entire telomerase is shown behind **c** and **f**.



**Fig. 2 |. TPP1 interactions with TERT TEN-TRAP.**

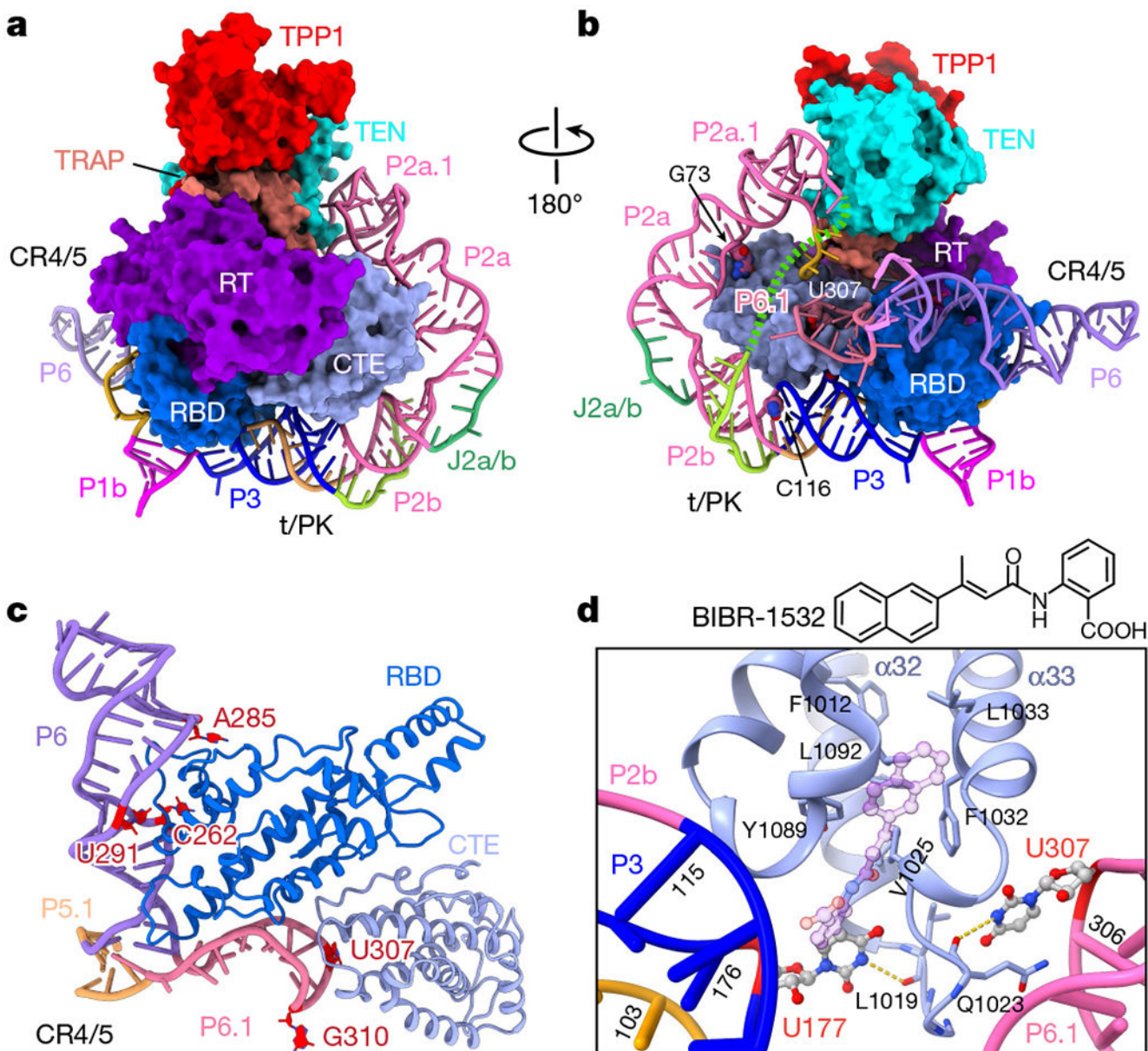
**a.** Cryo-EM maps of the catalytic core with (top) and without (bottom) TPP1. Both maps are low-pass filtered to 5 Å for comparison. **b.** Structure of TEN-TRAP with the extended  $\beta$ -sheet highlighted. **c.** Comparison of TPP1 structures in apo (gray) (Protein Data Bank (PDB): 2I46) and telomerase-bound form (red). NOB and TEL patch residues are in green and yellow, respectively. **d-f.** TPP1 interface with TEN-TRAP. **e.** TPP1 (red) sits on the top of TEN (cyan) and TRAP (salmon), with residues within 4 Å of the interface shown as spheres. **d, f.** Magnified view showing the polar (**d**) and hydrophobic (**f**) interactions. Boxed residues are positions with disease mutations. **g.** Effect of human TERT (hTERT) residue substitutions on telomerase activity stimulation by TPP1. Activity stimulation is calculated as activity with TPP1/activity without TPP1. Data are mean  $\pm$  s.d. from  $n = 3$  biologically independent experiments. \*\*\* $P < 0.001$ , \*\*\*\* $P < 0.0001$ ; one-tailed unpaired  $t$ -tests.





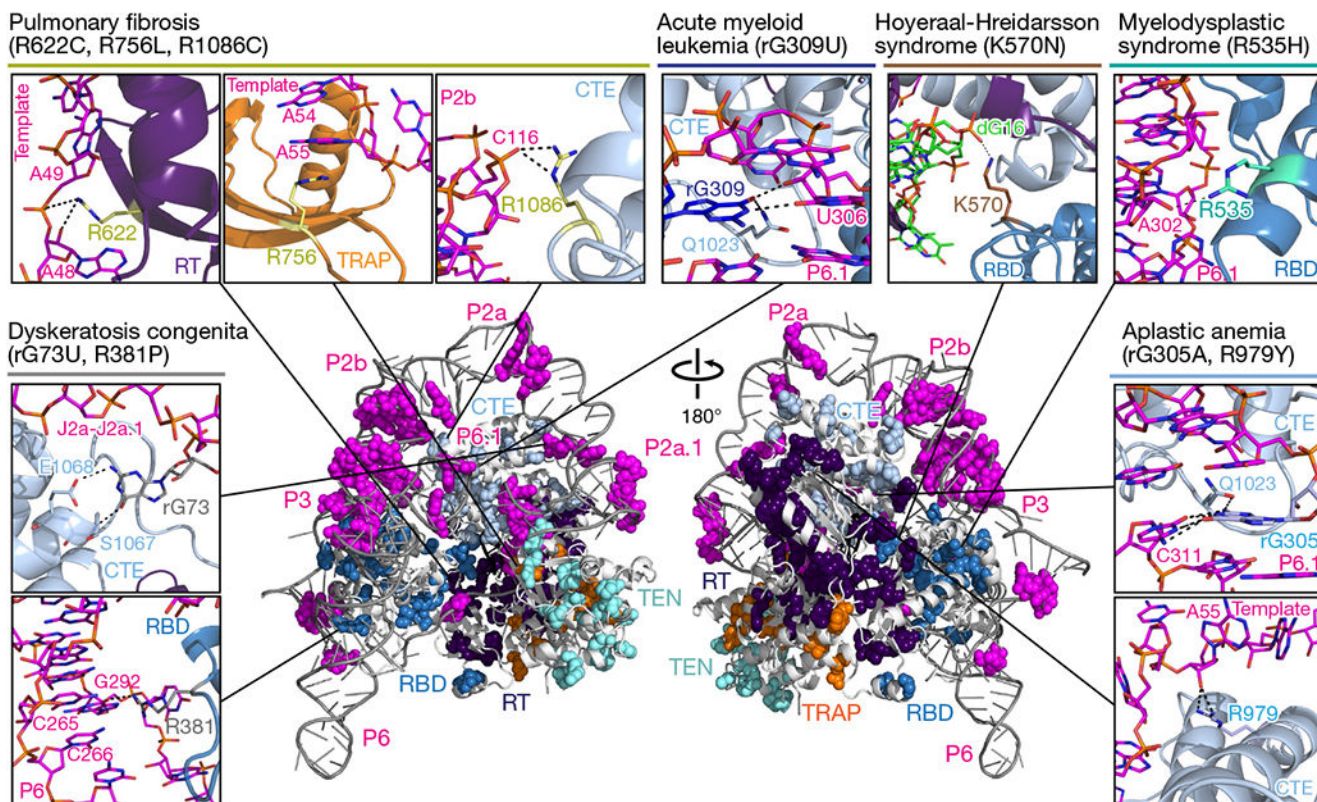
**Fig. 3 | Interactions between TERT and template–DNA duplex.**

**a, b,** The catalytic cavity of TERT (coloured surface) with TER t/PK (grey ribbon) and DNA (green ribbon). TERT is omitted in **b** to show the template–DNA duplex. The template and adjacent single-stranded regions are coloured red and orange, respectively. Top, template alignment nucleotides are shown in red with a white background. P1b and P2a.1 are shown in magenta. The red star represents the TERT active site. **c,** Detailed TERT–TER interactions in the region on the 5' side of the template. **d,** Ribbon depictions of template–DNA and TERT motifs involved in duplex handling. Bridge loop K499 and H500 are shown as sticks. **e,** Schematic showing specific interactions between TERT and template–DNA duplex. Polar and stacking interactions are indicated with arrows and bold lines, respectively. **f,** The path of TER nucleotides on the 3' side of the template. Positively charged TERT residues along the template exit path are shown as sticks.



**Fig. 4 | Interactions between TERT and TER.**

**a, b,** Telomerase catalytic core structure shown in two views. View in **a** is the same as Fig. 1b. The invisible linker between P2a.1 and the P2b loop is shown as a green dashed line in **b**. **c,** Structure of CR4/5 and its interactions with TERT RBD and CTE. Flipped out nucleotides (red) on P6 and P6.1 that interact with RBD and CTE, respectively, are labelled. G310 interacts with H2A–H2B. **d,** Top, the chemical structure of BIBR1532. Bottom, interactions of U177 (pseudoknot) and U307 (CR4/5) with CTE L<sub>α32-α33</sub> residues L1019 and Q1023, respectively. The telomerase-specific inhibitor BIBR1532 (light purple) is shown docked onto this structure based on the *T. castaneum* TERT-like protein CTE-BIBR1532 structure (PDB: 5CQG). Residues comprising the FVYL pocket for BIBR1532 binding<sup>11</sup> are labelled.



**Fig. 5 | Disease-associated mutations of human TERT and TER.**

Main image, cartoon representation of human TERT and TER (grey) with mutations associated with disease<sup>13</sup> (Supplementary Table 1) highlighted as spheres and coloured by their domains as in Fig. 1. Insets, magnified views of wild-type interactions in regions harbouring disease-causing mutations.